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FILE COVERS 1907 - 3 Feb 2003 VOL 138 ISS 6

FILE LAST UPDATED: 2 Feb 2003 (20030202/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

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L1	8	SEA FILE=REGISTRY	ABB=ON	PLU=ON	CEFTIOFUR/BI
L2	4	SEA FILE=REGISTRY	ABB=ON	PLU=ON	CEFIXIME/BI
L3	6	SEA FILE=REGISTRY	ABB=ON	PLU=ON	CEFOPERAZONE/BI
L4	26	SEA FILE=REGISTRY	ABB=ON	PLU=ON	CEFOTAXIME/BI
L5	9	SEA FILE=REGISTRY	ABB=ON	PLU=ON	CEFPODOXIME/BI
L6	8	SEA FILE=REGISTRY	ABB=ON	PLU=ON	CEFTAZIDIME/BI
L7	8	SEA FILE=REGISTRY	ABB=ON	PLU=ON	CEFTAZIDIME/BI
L8	4	SEA FILE=REGISTRY	ABB=ON	PLU=ON	CEFTIZOXIME/BI
L9	6	SEA FILE=REGISTRY	ABB=ON	PLU=ON	CEFTRIAXONE/BI
L10	3	SEA FILE=REGISTRY	ABB=ON	PLU=ON	CEFPPIROME/BI
L11	1	SEA FILE=REGISTRY	ABB=ON	PLU=ON	CEFLIDIN/BI
L12	6	SEA FILE=REGISTRY	ABB=ON	PLU=ON	CEFMENOXIME/BI
L13	3	SEA FILE=REGISTRY	ABB=ON	PLU=ON	CEFOZOPRAN/BI
L14	11614	SEA FILE=HCAPLUS	ABB=ON	PLU=ON	L1 OR L2 OR L3 OR L4 OR L5 OR L6 OR L7 OR L8 OR L9 OR L10 OR L11 OR L12 OR L13 OR CEFTIOFUR? OR CEFIPIME OR CEFIXIME OR CEFOPERAZONE OR CEFOTAXIME OR CEFPODOXIME
L15	6834	SEA FILE=HCAPLUS	ABB=ON	PLU=ON	CEFTAZIDIME OR CEFTAZIDIME OR CEFTIZOXIME OR CEFTRAXONE OR CEFTRIAXONE OR CEFPIROME OR CEFPPIROME OR CEFLIDIN OR CEFMENOXIME OR CEFOZOPRANE
L17 (	1048)	SEA FILE=REGISTRY	ABB=ON	PLU=ON	XYLAN/BI
L18 (	1750)	SEA FILE=REGISTRY	ABB=ON	PLU=ON	CHITIN/BI
L19 (	1443)	SEA FILE=REGISTRY	ABB=ON	PLU=ON	CHITOSAN/BI
L20 (	368)	SEA FILE=REGISTRY	ABB=ON	PLU=ON	CHONDROITIN/BI
L22 (	293)	SEA FILE=REGISTRY	ABB=ON	PLU=ON	ALGINATE/BI
L24 (	957)	SEA FILE=REGISTRY	ABB=ON	PLU=ON	PECTIN/BI
L25 (	1438)	SEA FILE=REGISTRY	ABB=ON	PLU=ON	(POLYSACCHARIDE/BI OR POLYSACCHARIDES/BI)
L28 (	1)	SEA FILE=REGISTRY	ABB=ON	PLU=ON	CYCLODEXTRINS/BI
L29 (	6)	SEA FILE=REGISTRY	ABB=ON	PLU=ON	CYCLOAMYLOSE/BI
L30 (	117)	SEA FILE=REGISTRY	ABB=ON	PLU=ON	CLATHRATE/BI
L32 (	2)	SEA FILE=REGISTRY	ABB=ON	PLU=ON	LIPOSOME/BI
L33 (	1)	SEA FILE=REGISTRY	ABB=ON	PLU=ON	POLYLACTIC (W) ACID
L52	64	SEA FILE=REGISTRY	ABB=ON	PLU=ON	BIOPOLYMER/BI
L53	30404	SEA FILE=HCAPLUS	ABB=ON	PLU=ON	L52 OR BIOPOLYMER
L54	231	SEA FILE=REGISTRY	ABB=ON	PLU=ON	(CARRAGEEN/BI OR CARRAGEENAN/ BI)
L55	10751	SEA FILE=HCAPLUS	ABB=ON	PLU=ON	L54 OR CARRAGEENAN

L56 28 SEA FILE=REGISTRY ABB=ON PLU=ON CARBOXYMETHYLCELLULOSE/BI  
 L57 6923 SEA FILE=HCAPLUS ABB=ON PLU=ON L56 OR CARBOXYMETHYLCELLULOSE  
  
 L58 516 SEA FILE=REGISTRY ABB=ON PLU=ON POLYPROPYLENE GLYCOL?/CN  
 L59 1778 SEA FILE=REGISTRY ABB=ON PLU=ON POLYETHYLENE GLYCOL?/CN  
 L60 170552 SEA FILE=HCAPLUS ABB=ON PLU=ON L59 OR (POLYETHYLENE OR POLY  
 (W) ETHYLENE) (2A) GLYCOL  
 L61 35804 SEA FILE=HCAPLUS ABB=ON PLU=ON L58 OR (POLYPROPYLENE OR POLY  
 (W) PROPYLENE) (2A) GLYCOL  
 L62 5 SEA FILE=REGISTRY ABB=ON PLU=ON POLYACETATE/BI  
 L63 404 SEA FILE=HCAPLUS ABB=ON PLU=ON L62 OR (POLYACETATE OR POLY  
 (W) ACETATE)  
 L64 39550 SEA FILE=HCAPLUS ABB=ON PLU=ON L32 OR LIPOSOME  
 L65 22952 SEA FILE=HCAPLUS ABB=ON PLU=ON L28 OR CYCLODEXTRIN  
 L66 166 SEA FILE=HCAPLUS ABB=ON PLU=ON L29 OR CYCLOAMYLOSE  
 L67 5668 SEA FILE=HCAPLUS ABB=ON PLU=ON L30 OR CLATHRATE  
 L68 9882 SEA FILE=HCAPLUS ABB=ON PLU=ON L29 OR AMYLOSE  
 L69 23 SEA FILE=REGISTRY ABB=ON PLU=ON POLY (L)XYLOSE  
 L70 51 SEA FILE=HCAPLUS ABB=ON PLU=ON L69 OR (POLYXYLOSE OR POLY  
 (2A) XYLOSE)  
 L71 4466 SEA FILE=HCAPLUS ABB=ON PLU=ON L33 OR (POLYLACTIC OR POLY  
 (W) LACTIC) (2A) ACID  
 L72 5923 SEA FILE=HCAPLUS ABB=ON PLU=ON L17 OR XYLAN  
 L73 11676 SEA FILE=HCAPLUS ABB=ON PLU=ON L18 OR CHITIN  
 L74 13648 SEA FILE=HCAPLUS ABB=ON PLU=ON L19 OR CHITOSAN  
 L75 12206 SEA FILE=HCAPLUS ABB=ON PLU=ON L20 OR CHONDROITIN (W)  
 SULFATE  
 L76 18083 SEA FILE=HCAPLUS ABB=ON PLU=ON L22 OR (SODIUM OR NA) (W)  
 ALGINAT?  
 L77 19827 SEA FILE=HCAPLUS ABB=ON PLU=ON L24 OR PECTIN  
 L78 74282 SEA FILE=HCAPLUS ABB=ON PLU=ON L25 OR POLYSACCHARIDE  
 L79 332 SEA FILE=HCAPLUS ABB=ON PLU=ON FATTY (W) ACID (W) COMPLEX  
 L81 75 SEA FILE=HCAPLUS ABB=ON PLU=ON (L14 OR L15) (L) (L53 OR L55  
 OR L72 OR L73 OR L74 OR L75 OR L76 OR L57 OR L77 OR L78 OR L61  
 OR L60 OR L63 OR L64 OR L79 OR L65 OR L66 OR L67 OR L68 OR L70  
 OR L71)  
 L82 73448 SEA FILE=REGISTRY ABB=ON PLU=ON CALCIUM  
 L83 78644 SEA FILE=REGISTRY ABB=ON PLU=ON MAGNESIUM  
 L84 86473 SEA FILE=REGISTRY ABB=ON PLU=ON LITHIUM  
 L85 1 SEA FILE=REGISTRY ABB=ON PLU=ON IRON/CN  
 L86 1 SEA FILE=REGISTRY ABB=ON PLU=ON COPPER/CN  
 L87 129382 SEA FILE=REGISTRY ABB=ON PLU=ON ZINC  
 L88 197971 SEA FILE=REGISTRY ABB=ON PLU=ON ALUMINUM  
 L89 142592 SEA FILE=REGISTRY ABB=ON PLU=ON MANGANESE  
 L90 7 SEA FILE=HCAPLUS ABB=ON PLU=ON L81 AND (CALCIUM OR L82 OR  
 L83 OR L84 OR L85 OR L86 OR L87 OR L88 OR L89 OR MAGNESIUM OR  
 LITHIUM OR IRON OR COPPER OR ZINC OR ALUMINUM OR MANGANESE)

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L90 ANSWER 1 OF 7 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER: 2002:675842 HCAPLUS

DOCUMENT NUMBER: 137:206570

TITLE: Oral pharmaceutical composition of cefpodoxime  
proxetil

INVENTOR(S): Malhotra, Mukta; Mathur, Rajeev Shankar; Malik, Rajiv

PATENT ASSIGNEE(S): Ranbaxy Laboratories Limited, India

SOURCE: PCT Int. Appl., 20 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

## PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002067943	A1	20020906	WO 2002-IB602	20020227
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
PRIORITY APPLN. INFO.:			IN 2001-DE190	A 20010227
AB The present invention relates to a stable pharmaceutical composition of cefpodoxime proxetil (I), for oral administration. A tablet contained I 49.79, <b>calcium</b> CM-cellulose 42.07, lactose 3.64, hydroxypropyl cellulose 1.64, sodium lauryl sulfate 2.05, and <b>magnesium</b> stearate 0.82%. The amount of I released in glycine buffer after 10 min was 88.8%.				
IT <b>9000-07-1, Carrageenan gum 9003-11-6, Polyoxyethylene polyoxypropylene copolymer 9050-04-8, Calcium</b> carboxymethyl cellulose <b>11138-66-2</b> , Xanthan gum RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (oral pharmaceutical composition of <b>cefpodoxime</b> proxetil)				
REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT				
L90 ANSWER 2 OF 7 HCAPLUS COPYRIGHT 2003 ACS				
ACCESSION NUMBER: 2001:935443 HCAPLUS				
DOCUMENT NUMBER: 136:58849				
TITLE: Compositions and methods to improve the oral absorption of antimicrobial agents				
INVENTOR(S): Choi, Seung-Ho; Lee, Jeoung-Soo; Keith, Dennis				
PATENT ASSIGNEE(S): Cubist Pharmaceuticals, Inc., USA; International Health Management Associates, Inc.; University of Utah Research Foundation				
SOURCE: PCT Int. Appl., 70 pp. CODEN: PIXXD2				
DOCUMENT TYPE: Patent				
LANGUAGE: English				
FAMILY ACC. NUM. COUNT: 2				
PATENT INFORMATION:				

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2001097851	A2	20011227	WO 2001-US19625	20010618
WO 2001097851	A3	20020516		
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG				
US 6248360	B1	20010619	US 2000-598089	20000621
PRIORITY APPLN. INFO.:			US 2000-598089	A 20000621
			US 2001-829405	A 20010409
			US 2001-283976P	P 20010416

AB The present invention provides compns. and methods for increasing absorption of antibacterial agents, particularly third generation cephalosporin antibacterial agents, in oral dosage solid and/or suspension forms. Specifically, the composition is comprised of a **biopolymer** that is preferably swellable and/or mucoadhesive, an antimicrobial agent, and a cationic binding agent contained within the **biopolymer** such that the binding agent is ionically bound or complexed to at least one member selected from the group consisting of the **biopolymer** and the antimicrobial agent. A solution of 44.5 mg calcium chloride in 10 mL water and 1.0 g of **ceftriaxone** in 10 mL water was added gradually to a solution of 400 mg **carrageenan** and the dispersion was centrifuged and the supernatant was lyophilized. The resulting composition comprized **carrageenan** 27.7, **ceftriaxone** 1, and **calcium** chloride 3.1%. Plasma concentration of different antimicrobial-**biopolymer** complexes after oral administration to rats was measured.

IT 7429-90-5DP, **Aluminum**, conjugates with biopolymers and antimicrobial agents 7439-89-6DP, **Iron**, conjugates with biopolymers and antimicrobial agents 7439-93-2DP, **Lithium**, conjugates with biopolymers and antimicrobial agents 7439-95-4DP, **Magnesium**, conjugates with biopolymers and antimicrobial agents 7439-96-5DP, **Manganese**, conjugates with biopolymers and antimicrobial agents 7440-50-8DP, **Copper**, conjugates with biopolymers and antimicrobial agents 7440-66-6DP, **Zinc**, conjugates with biopolymers and antimicrobial agents 7440-70-2DP, **Calcium**, conjugates with biopolymers and antimicrobial agents 10043-52-4DP, **Calcium** chloride, conjugates with antimicrobials and biopolymers 62893-19-0DP, **Cefoperazone**, conjugates with biopolymers and cationic binding agents 63527-52-6DP, **Cefotaxime**, conjugates with biopolymers and cationic binding agents 65085-01-0DP, **Cefmenoxime**, conjugates with biopolymers and cationic binding agents 68401-81-0DP, **Ceftizoxime**, conjugates with biopolymers and cationic binding agents 72558-82-8DP, **Ceftazidime**, conjugates with biopolymers and cationic binding agents 73384-59-5DP, **Ceftriaxone**, conjugates with biopolymers and cationic binding agents 79350-37-1DP, **Cefixime**, conjugates with biopolymers and cationic binding agents 80210-62-4DP, **Cefpodoxime**, conjugates with biopolymers and cationic binding agents 80370-57-6DP, **Ceftiofur**, conjugates with biopolymers and cationic binding agents 84957-29-9DP, **Cefpirome**, conjugates with biopolymers and cationic binding agents 105239-91-6DP, **Cefclidin**, conjugates with biopolymers and cationic binding agents 113359-04-9DP, **Cefozopran**, conjugates with biopolymers and cationic binding agents  
 RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(compns. and methods to improve oral absorption of antimicrobial agents)

L90 ANSWER 3 OF 7 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER: 2001:449182 HCAPLUS

DOCUMENT NUMBER: 135:51066

TITLE: Complexes to improve oral absorption of poorly absorbable antibiotics

INVENTOR(S): Choi, Seung-ho; Lee, Jeoung-soo

PATENT ASSIGNEE(S): International Health Management Associates, Inc., USA

SOURCE: U.S., 9 pp.

CODEN: USXXAM

DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 2  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 6248360	B1	20010619	US 2000-598089	20000621
WO 2001097851	A2	20011227	WO 2001-US19625	20010618
WO 2001097851	A3	20020516		

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM  
 RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

PRIORITY APPLN. INFO.:  
 US 2000-598089 A 20000621  
 US 2001-829405 A 20010409  
 US 2001-283976P P 20010416

AB The present invention provides compns. and methods for increasing absorption of poorly absorbable antibiotics, particularly third generation cephalosporin antibiotics, in oral dosage solid and/or suspension forms. Specifically, the composition is comprised of a **biopolymer** that is preferably swellable and/or mucoadhesive, a poorly absorbable antibiotic, and a cationic binding agent contained within the **biopolymer** such that the binding agent is tonically bound or complexed to at least 1 member selected from the group consisting of the **biopolymer** and the antibiotic. A **ceftriaxone-carrageenan-calcium** complex was prepared by the treatment of the antibiotic with **calcium** and **carrageenan**. The plasma drug concentration from the complex was greater than that obtained by administering the antibiotic in an uncomplexed state.

IT 7440-66-6DP, Zinc, antibiotic complexes, biological studies 7440-70-2DP, Calcium, antibiotic complexes, biological studies 73384-59-5DP, Ceftriaxone, carrageenan complexes

RL: BPR (Biological process); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses)

(complexes for improvement of oral absorption of poorly absorbable antibiotics)

IT 7429-90-5, Aluminum, biological studies  
 7439-89-6, Iron, biological studies 7439-95-4, Magnesium, biological studies 7439-96-5, Manganese, biological studies 7440-50-8, Copper, biological studies

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(complexes for improvement of oral absorption of poorly absorbable antibiotics)

REFERENCE COUNT: 8 THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L90 ANSWER 4 OF 7 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER: 2001:231880 HCAPLUS

DOCUMENT NUMBER: 136:11010

TITLE: Gel formation of cefpodoxime proxetil, basic antibiotic and its formulation design

AUTHOR(S): Hamaura, Takeshi

CORPORATE SOURCE: Product Development Laboratories, Sankyo Co., Ltd., Hiromachi, Shinagawa-ku, Tokyo, Japan

SOURCE: Pharm Tech Japan (2001), 17(4), 619-624, 627-632  
CODEN: PTJAE9; ISSN: 0910-4739  
PUBLISHER: Jiho  
DOCUMENT TYPE: Journal  
LANGUAGE: Japanese

AB Cefpodoxime proxetil (CPDX-PR), an orally-active cephalosporin antibiotic, possesses one asym. carbon atom in the ester group and exists as a mixture of two amorphous diastereoisomers, CPDX-PR (A) and CPDX-PR (B). There are marked differences in the dissoln. behavior of CPDX-PRs (A) and (B). Although the dissolved amount of CPDX-PR(A) increases with time to reach equilibrium rapidly, CPDX-PR (B) exhibits supersatd. phenomena in strongly-acidic conditions. Super-saturated solns. of CPDX-PR(B) remain as clear solns. at first, but finally transform into gels. From elec. microscope data, it was revealed that spherical particles of crystalline CPDX-PR(B) HCl salt adhere together to constitute the framework of the gel and solution occupies gaps in the framework of the gel. CPDX-PR model tablets formulated a disintegrant with comparably low swelling pressure, exhibited decreased dissoln. in strongly-acidic conditions. A gel layer formed at the surface of the tablets must inhibit water penetration into the inner part of the tablets. Thus, the tablets disintegrate slowly, resulting in poor dissoln. However, formulation of a disintegrant having excellent swelling behavior was revealed to improve both the delayed disintegration and the decreased dissoln. of CPDX-PR tablets. The tablets with the improved dissoln. behavior apparently exhibited higher in vivo bioavailability in beagle dogs than that with the decreased dissoln. due to gel formation.

IT 9050-04-8

RL: MOA (Modifier or additive use); THU (Therapeutic use); BIOL  
(Biological study); USES (Uses)  
(gel formation of **cefpodoxime proxetil** and its formulation design)

L90 ANSWER 5 OF 7 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER: 1997:795285 HCAPLUS  
DOCUMENT NUMBER: 128:110395  
TITLE: Compatibility of doxorubicin hydrochloride liposome injection with selected other drugs during simulated Y-site administration  
AUTHOR(S): Trissel, Lawrence A.; Gilbert, Doward L.; Martinez, Juan F.  
CORPORATE SOURCE: Division of Pharmacy, The University of Texas M. D. Anderson Cancer Center, Houston, TX, 77030, USA  
SOURCE: American Journal of Health-System Pharmacy (1997), 54(23), 2708-2713  
CODEN: AHSPEK; ISSN: 1079-2082  
PUBLISHER: American Society of Health-System Pharmacists  
DOCUMENT TYPE: Journal  
LANGUAGE: English

AB The compatibility of doxorubicin hydrochloride liposome injection with selected other drugs during simulated Y-site administration was studied. Five milliliters of doxorubicin hydrochloride liposome injection 0.4 mg/mL in 5% dextrose injection was combined with 5 mL of each of 82 other drugs in 5% dextrose injection or, if necessary to avoid incompatibilities with the diluent, 0.9% sodium chloride injection. The combinations were examined with the unaided eye in fluorescent light and in high-intensity monodirectional light to enhance visualization of small particles and low-level turbidity. The turbidity of each combination was measured as well. Particle sizing and counting were performed on selected combinations. Evaluations were performed initially and at one and four hours. All combinations were stored at room temperature (.apprx.23 °C). Most of the test drugs were compatible with doxorubicin hydrochloride liposome injection during the four-hour observation period. However, practitioners should be cautious in administering any drug simultaneously

with doxorubicin hydrochloride liposome injection until the integrity of the liposomes can be verified. Eighteen drugs exhibited unacceptable increases or decreases in measured turbidity or particulate formation within four hours. During simulated Y-site administration, doxorubicin hydrochloride 0.4 mg/mL (as the liposomal injection) in 5% dextrose injection was compatible with 64 of 82 other drugs for four hours at .apprx.23 °C and was incompatible with 18 of the test drugs.

IT 299-28-5, Calcium gluconate 1492-18-8,  
Leucovorin calcium 7487-88-9, Magnesium  
sulfate, biological studies 62893-20-3, Cefoperazone  
sodium 68401-82-1, Ceftizoxime sodium  
73547-61-2, Ceftazidime sodium 74578-69-1,  
Ceftriaxone sodium  
RL: ADV (Adverse effect, including toxicity); BIOL (Biological study)  
(doxorubicin hydrochloride liposome injection compatibility  
with other drugs)

L90 ANSWER 6 OF 7 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER: 1995:1003883 HCAPLUS  
DOCUMENT NUMBER: 124:97432  
TITLE: Decrease in dissolution of cefpodoxime proxetil  
tablets by gel formation and its improvement  
AUTHOR(S): Hamaura, Takeshi; Terashima, Hidenari; Ohtani, Tomoko;  
Mori, Yohko; Seta, Yasuo; Kunihiro, Kusai; Kenji,  
Sasahara; Nishimura, Kenji  
CORPORATE SOURCE: Product Development Laboratories, Sankyo Company Ltd.,  
Tokyo, 140, Japan  
SOURCE: Yakuzai (1995), 55(3), 175-82  
CODEN: YAKUA2; ISSN: 0372-7629  
PUBLISHER: Nippon Yakuzai Gakkai  
DOCUMENT TYPE: Journal  
LANGUAGE: Japanese

AB Cefpodoxime proxetil (CPDX-PR) has been developed as a racemate and only one of the two diastereoisomers forms a gel in strongly acidic conditions. CPDX-PR model tablets, formulated with a disintegrant with low swelling pressure, exhibited decreased dissoln. in strongly acetic conditions. These phenomena were explained by formation of a gel layer at tablet surface, inhibiting water penetration into the inner part of the tablets. Thus, the tablets are slow to disintegrate, resulting in poor dissoln. However, formulation with a disintegrant having excellent swelling behavior improved both the delayed disintegration and the decreased dissoln. of CPDX-PR tablets. Tablets with the improved dissoln. behavior apparently exhibited higher in vivo bioavailability in beagle dogs than those with the decreased dissoln. due to gel formation.

IT 9050-04-8, CM-cellulose calcium  
RL: MOA (Modifier or additive use); THU (Therapeutic use); BIOL  
(Biological study); USES (Uses)  
(disintegrant; decrease in dissoln. of cefpodoxime proxetil  
tablets by gel formation and its improvement)

L90 ANSWER 7 OF 7 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER: 1991:192593 HCAPLUS  
DOCUMENT NUMBER: 114:192593  
TITLE: Nonphospholipid pharmaceutical liposomes  
INVENTOR(S): Radhakrishnan, Ramachandran  
PATENT ASSIGNEE(S): Liposome Technology, Inc., USA  
SOURCE: PCT Int. Appl., 96 pp.  
CODEN: PIXXD2  
DOCUMENT TYPE: Patent  
LANGUAGE: English  
FAMILY ACC. NUM. COUNT: 2  
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9006775	A1	19900628	WO 1989-US5525	19891206
W: AU, DK, FI, JP, NO				
RW: AT, BE, CH, DE, ES, FR, GB, IT, LU, NL, SE				
US 4906476	A	19900306	US 1988-284158	19881214
US 5043165	A	19910827	US 1988-284216	19881214
PRIORITY APPLN. INFO.:			US 1988-284158	19881214
			US 1988-284216	19881214

AB A nonconventional liposome composition consisting of nonphospholipid lipids, especially cholesterol and cholesterol ester salts, are used for encapsulation of drugs. They are useful for sustained release of steroids, and are suitable for treatment of inflammatory, arthritic, rheumatoid diseases, etc., especially as aerosols for interstitial lung disease. Beclomethasone dipropionate (I) 10 was incorporated into liposomes prepared with Na cholesterol sulfate 50 and cholesterol 40 mol %. Sustained release of I was observed in rats following intratracheal administration, in contrast to liposomes formulated with phosphatidylcholine and cholesterol.

IT 24352-55-4 99523-97-4 133058-04-5  
133058-05-6 133161-25-8 133161-26-9  
133352-85-9 133352-86-0 133442-38-3

RL: BIOL (Biological study)  
(pharmaceutical liposomes containing cholesterol and)

IT 63527-52-6, Cefotaxime

RL: BIOL (Biological study)  
(pharmaceutical liposomes containing cholesterol and salt of cholesterol esterand)

=> => d stat que

L1	8	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFTIOFUR/BI
L2	4	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFIXIME/BI
L3	6	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFOPERAZONE/BI
L4	26	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFOTAXIME/BI
L5	9	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFPDOXIME/BI
L6	8	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFTAZIDIME/BI
L7	8	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFTAZIDIME/BI
L8	4	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFTIZOXIME/BI
L9	6	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFTRIAXONE/BI
L10	3	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFPPIROME/BI
L11	1	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFCLIDIN/BI
L12	6	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFMENOXIME/BI
L13	3	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFOZOPRAN/BI
L14	11614	SEA	FILE=HCAPLUS	ABB=ON	PLU=ON	L1 OR L2 OR L3 OR L4 OR L5 OR L6 OR L7 OR L8 OR L9 OR L10 OR L11 OR L12 OR L13 OR CEFTIOFUR? OR CEFIPIME OR CEFIXIME OR CEFOPERAZONE OR CEFOTAXIME OR CEFPDOXIME
L15	6834	SEA	FILE=HCAPLUS	ABB=ON	PLU=ON	CEFTAZIDIME OR CEFTAZIDIME OR CEFTIZOXIME OR CEFTRAXONE OR CEFTRIAXONE OR CEFPIROME OR CEFCLIDIN OR CEFMENOXIME OR CEFOZOPRANE
L17 (	1048)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	XYLAN/BI
L18 (	1750)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CHITIN/BI
L19 (	1443)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CHITOSAN/BI
L20 (	368)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CHONDROITIN/BI
L22 (	293)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	ALGINATE/BI
L24 (	957)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	PECTIN/BI
L25 (	1438)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	(POLYSACCHARIDE/BI OR POLYSACCHARIDES/BI)
L28 (	1)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CYCLODEXTRINS/BI
L29 (	6)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CYCLOAMYLOSE/BI
L30 (	117)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CLATHRATE/BI
L32 (	2)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	LIPOSOME/BI
L33 (	1)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	POLYLACTIC (W) ACID



L52	64	SEA FILE=REGISTRY ABB=ON	PLU=ON	BIOPOLYMER/BI
L53	30404	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L52 OR BIOPOLYMER
L54	231	SEA FILE=REGISTRY ABB=ON	PLU=ON	(CARRAGEEN/BI OR CARRAGEENAN/BI)
L55	10751	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L54 OR CARRAGEENAN
L56	28	SEA FILE=REGISTRY ABB=ON	PLU=ON	CARBOXYMETHYLCELLULOSE/BI
L57	6923	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L56 OR CARBOXYMETHYLCELLULOSE
L58	516	SEA FILE=REGISTRY ABB=ON	PLU=ON	POLYPROPYLENE GLYCOL?/CN
L59	1778	SEA FILE=REGISTRY ABB=ON	PLU=ON	POLYETHYLENE GLYCOL?/CN
L60	170552	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L59 OR (POLYETHYLENE OR POLY(W) ETHYLENE) (2A) GLYCOL
L61	35804	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L58 OR (POLYPROPYLENE OR POLY(W) PROPYLENE) (2A) GLYCOL
L62	5	SEA FILE=REGISTRY ABB=ON	PLU=ON	POLYACETATE/BI
L63	404	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L62 OR (POLYACETATE OR POLY(W) ACETATE)
L64	39550	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L32 OR LIPOSOME
L65	22952	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L28 OR CYCLODEXTRIN
L66	166	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L29 OR CYCLOAMYLOSE
L67	5668	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L30 OR CLATHRATE
L68	9882	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L29 OR AMYLOSE
L69	23	SEA FILE=REGISTRY ABB=ON	PLU=ON	POLY (L)XYLOSE
L70	51	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L69 OR (POLYXYLOSE OR POLY(2A) XYLOSE)
L71	4466	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L33 OR (POLYLACTIC OR POLY(W) LACTIC) (2A) ACID
L72	5923	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L17 OR XYLAN
L73	11676	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L18 OR CHITIN
L74	13648	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L19 OR CHITOSAN
L75	12206	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L20 OR CHONDROITIN (W) SULFATE
L76	18083	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L22 OR (SODIUM OR NA) (W) ALGINAT?
L77	19827	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L24 OR PECTIN
L78	74282	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L25 OR POLYSACCHARIDE
L79	332	SEA FILE=HCAPLUS ABB=ON	PLU=ON	FATTY (W) ACID (W) COMPLEX
L81	75	SEA FILE=HCAPLUS ABB=ON	PLU=ON	(L14 OR L15) (L) (L53 OR L55 OR L72 OR L73 OR L74 OR L75 OR L76 OR L57 OR L77 OR L78 OR L61 OR L60 OR L63 OR L64 OR L79 OR L65 OR L66 OR L67 OR L68 OR L70 OR L71)
L82	73448	SEA FILE=REGISTRY ABB=ON	PLU=ON	CALCIUM
L83	78644	SEA FILE=REGISTRY ABB=ON	PLU=ON	MAGNESIUM
L84	86473	SEA FILE=REGISTRY ABB=ON	PLU=ON	LITHIUM
L85	1	SEA FILE=REGISTRY ABB=ON	PLU=ON	IRON/CN
L86	1	SEA FILE=REGISTRY ABB=ON	PLU=ON	COPPER/CN
L87	129382	SEA FILE=REGISTRY ABB=ON	PLU=ON	ZINC
L88	197971	SEA FILE=REGISTRY ABB=ON	PLU=ON	ALUMINUM
L89	142592	SEA FILE=REGISTRY ABB=ON	PLU=ON	MANGANESE
L90	7	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L81 AND (CALCIUM OR L82 OR L83 OR L84 OR L85 OR L86 OR L87 OR L88 OR L89 OR MAGNESIUM OR LITHIUM OR IRON OR COPPER OR ZINC OR ALUMINUM OR MANGANESE)
L91	1	SEA FILE=HCAPLUS ABB=ON	PLU=ON	(L81 AND METAL) NOT L90

=> d ibib abs hitrn l91 tot

L91 ANSWER 1 OF 1 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER: 1999:8553 HCAPLUS

DOCUMENT NUMBER: 130:177099

TITLE: Studies on drugs absorption using liposome to simulate biomembrane

AUTHOR(S): Luo, Yun-Jing; Shen, Han-Xi; Liu, Hai-Tao; Kong,

De-Ming  
 CORPORATE SOURCE: Department of Chemistry, Nankai University, Tianjin,  
 300071, Peop. Rep. China  
 SOURCE: Gaodeng Xuexiao Huaxue Xuebao (1998), 19(11),  
 1730-1734  
 CODEN: KTHPDM; ISSN: 0251-0790  
 PUBLISHER: Gaodeng Jiaoyu Chubanshe  
 DOCUMENT TYPE: Journal  
 LANGUAGE: Chinese

AB The absorptions of liposol. salicylic acid, water-soluble vitamin B6, **ceftriaxone**, the quinolone antibacterial norfloxacin and anticancer doxorubicin were discussed by using, **liposome** as biomembrane model and comparison with cell membrane. The absorption of salicylic acid whose absorption curve is linear depends on the passive diffusion. The exptl. result of vitamin B6 supports the opinion that the absorption of vitamin B6 depends on proteins which lie in membranes. The absorption of vitamin B6 is a chem. process, so absorption curve is nonlinear. The absorption mode of **ceftriaxone** is a simple diffusion. The absorption of norfloxacin depends on the effects of **metal** ions. Under the driving force of nucleic acid in cells, doxorubicin was absorbed through biomembrane.

IT 73384-59-5, **Ceftriaxone**  
 RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)  
 (studies on drugs absorption using **liposome** to simulate biomembrane)

=> => d stat que

L1	8	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFTIOFUR/BI
L2	4	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFIXIME/BI
L3	6	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFOPERAZONE/BI
L4	26	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFOTAXIME/BI
L5	9	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFPODOXIME/BI
L6	8	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFTAZIDIME/BI
L7	8	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFTAZIDIME/BI
L8	4	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFTIZOXIME/BI
L9	6	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFTRIAZONE/BI
L10	3	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFPRIOME/BI
L11	1	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFCLIDIN/BI
L12	6	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFMENOXIME/BI
L13	3	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CEFOZOPRAN/BI
L14	11614	SEA	FILE=HCAPLUS	ABB=ON	PLU=ON	L1 OR L2 OR L3 OR L4 OR L5 OR L6 OR L7 OR L8 OR L9 OR L10 OR L11 OR L12 OR L13 OR CEFTIOFUR? OR CEFPIPIE OR CEFIXIME OR CEFOPERAZONE OR CEFOTAXIME OR CEFPODOXIME
L15	6834	SEA	FILE=HCAPLUS	ABB=ON	PLU=ON	CEFTAZIDIME OR CEFTAZIDIME OR CEFTIZOXIME OR CEFTRAXONE OR CEFTRIAZONE OR CEFPRIOME OR CEFPIROME OR CEFCLIDIN OR CEFMENOXIME OR CEFOZOPRANE
L17 (	1048)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	XYLAN/BI
L18 (	1750)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CHITIN/BI
L19 (	1443)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CHITOSAN/BI
L20 (	368)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CHONDROITIN/BI
L22 (	293)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	ALGINATE/BI
L24 (	957)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	PECTIN/BI
L25 (	1438)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	(POLYSACCHARIDE/BI OR POLYSACCHARIDES/BI)
L28 (	1)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CYCLODEXTRINS/BI
L29 (	6)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CYCLOAMYLOSE/BI
L30 (	117)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	CLATHRATE/BI
L32 (	2)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	LIPOSOME/BI
L33 (	1)	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	POLYLACTIC (W) ACID
L52	64	SEA	FILE=REGISTRY	ABB=ON	PLU=ON	BIOPOLYMER/BI

L53	30404	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L52 OR BIOPOLYMER
L54	231	SEA FILE=REGISTRY ABB=ON	PLU=ON	(CARRAGEEN/BI OR CARRAGEENAN/BI)
L55	10751	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L54 OR CARRAGEENAN
L56	28	SEA FILE=REGISTRY ABB=ON	PLU=ON	CARBOXYMETHYLCELLULOSE/BI
L57	6923	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L56 OR CARBOXYMETHYLCELLULOSE
L58	516	SEA FILE=REGISTRY ABB=ON	PLU=ON	POLYPROPYLENE GLYCOL?/CN
L59	1778	SEA FILE=REGISTRY ABB=ON	PLU=ON	POLYETHYLENE GLYCOL?/CN
L60	170552	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L59 OR (POLYETHYLENE OR POLY(W) ETHYLENE) (2A) GLYCOL
L61	35804	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L58 OR (POLYPROPYLENE OR POLY(W) PROPYLENE) (2A) GLYCOL
L62	5	SEA FILE=REGISTRY ABB=ON	PLU=ON	POLYACETATE/BI
L63	404	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L62 OR (POLYACETATE OR POLY(W) ACETATE)
L64	39550	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L32 OR LIPOSOME
L65	22952	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L28 OR CYCLODEXTRIN
L66	166	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L29 OR CYCLOAMYLOSE
L67	5668	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L30 OR CLATHRATE
L68	9882	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L29 OR AMYLOSE
L69	23	SEA FILE=REGISTRY ABB=ON	PLU=ON	POLY (L)XYLOSE
L70	51	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L69 OR (POLYXYLOSE OR POLY(2A) XYLOSE)
L71	4466	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L33 OR (POLYLACTIC OR POLY(W) LACTIC) (2A) ACID
L72	5923	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L17 OR XYLAN
L73	11676	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L18 OR CHITIN
L74	13648	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L19 OR CHITOSAN
L75	12206	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L20 OR CHONDROITIN (W) SULFATE
L76	18083	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L22 OR (SODIUM OR NA) (W) ALGINAT?
L77	19827	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L24 OR PECTIN
L78	74282	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L25 OR POLYSACCHARIDE
L79	332	SEA FILE=HCAPLUS ABB=ON	PLU=ON	FATTY (W) ACID (W) COMPLEX
L81	75	SEA FILE=HCAPLUS ABB=ON	PLU=ON	(L14 OR L15) (L) (L53 OR L55 OR L72 OR L73 OR L74 OR L75 OR L76 OR L57 OR L77 OR L78 OR L61 OR L60 OR L63 OR L64 OR L79 OR L65 OR L66 OR L67 OR L68 OR L70 OR L71)
L82	73448	SEA FILE=REGISTRY ABB=ON	PLU=ON	CALCIUM
L83	78644	SEA FILE=REGISTRY ABB=ON	PLU=ON	MAGNESIUM
L84	86473	SEA FILE=REGISTRY ABB=ON	PLU=ON	LITHIUM
L85	1	SEA FILE=REGISTRY ABB=ON	PLU=ON	IRON/CN
L86	1	SEA FILE=REGISTRY ABB=ON	PLU=ON	COPPER/CN
L87	129382	SEA FILE=REGISTRY ABB=ON	PLU=ON	ZINC
L88	197971	SEA FILE=REGISTRY ABB=ON	PLU=ON	ALUMINUM
L89	142592	SEA FILE=REGISTRY ABB=ON	PLU=ON	MANGANESE
L90	7	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L81 AND (CALCIUM OR L82 OR L83 OR L84 OR L85 OR L86 OR L87 OR L88 OR L89 OR MAGNESIUM OR LITHIUM OR IRON OR COPPER OR ZINC OR ALUMINUM OR MANGANESE)
L91	1	SEA FILE=HCAPLUS ABB=ON	PLU=ON	(L81 AND METAL) NOT L90
L92	7	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L81 AND (ORAL? OR MOUTH)
L93	3	SEA FILE=HCAPLUS ABB=ON	PLU=ON	L92 NOT (L90 OR L91)

=> d ibib abs hitrn 193 tot

L93 ANSWER 1 OF 3 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER: 2002:391569 HCAPLUS

DOCUMENT NUMBER: 136:374901

TITLE: Antibiotics adducts with natural polysaccharide polymers in the form of aqueous solutions

INVENTOR(S): Anzaghi, Piergiorgio; Stefli, Rosanna  
 PATENT ASSIGNEE(S): Pharma Biotech Limited, Cyprus  
 SOURCE: PCT Int. Appl., 16 pp.  
 CODEN: PIXXD2  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002040057	A1	20020523	WO 2001-IB2172	20011116
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
AU 2002012632	A5	20020527	AU 2002-12632	20011116
PRIORITY APPLN. INFO.: IT 2000-MI2478 A 20001117 WO 2001-IB2172 W 20011116				
AB Antibiotics adducts with natural <b>polysaccharide</b> polymers in the form of aqueous solns., endowed with better antibacterial activity if compared to the starting antibiotic are described. An extemporaneous solution of the adduct 60% <b>ceftriaxone</b> -40% dextran 5 was prepared				
IT <b>73384-59-5DP, Ceftriaxone</b> , adducts with dextran RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (antibiotics adducts with natural <b>polysaccharide</b> polymers in the form of aqueous solns.)				

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS  
 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L93 ANSWER 2 OF 3 HCAPLUS COPYRIGHT 2003 ACS  
 ACCESSION NUMBER: 1998:261905 HCAPLUS  
 DOCUMENT NUMBER: 129:32211  
 TITLE: Cefpodoxime-proxetil protection from intestinal lumen  
 hydrolysis by oil-in-water submicron emulsions  
 AUTHOR(S): Crauste-Manciet, Sylvie; Brossard, Denis; Decroix,  
 Marie-Odile; Farinotti, Robert; Chaumeil, Jean-Claude  
 CORPORATE SOURCE: Laboratoire de Pharmacotechnie et Dermopharmacie,  
 Faculte des Sciences Pharmaceutiques et Biologiques,  
 Paris V, Paris, 75006, Fr.  
 SOURCE: International Journal of Pharmaceutics (1998), 165(1),  
 97-106  
 CODEN: IJPHDE; ISSN: 0378-5173  
 PUBLISHER: Elsevier Science B.V.  
 DOCUMENT TYPE: Journal  
 LANGUAGE: English  
 AB Cefpodoxime proxetil is an **orally** active, broad spectrum, third  
 generation cephalosporin ester. This prodrug was previously found to be  
 hydrolyzed in vitro both in rabbit and human duodenal washing by a  
 cholinesterase. The objective of this work was to find a formulation  
 which can protect the prodrug from enzymic attack. In order to protect  
 the prodrug from enzymic hydrolysis, the objective was to include it into  
 the oil phase of an oil-in-water (o/w) emulsion. Somehow, cefpodoxime  
 proxetil posed specific problems related to the solubilization. The  
 solubilization was obtained with a mixed medium-chain-triglycerides  
 (MCT)/blends of mono-, di- and triglycerides oil phase and the optimal

ratio was defined to be 60:40 (weight/weight) in order to obtain emulsification.

The emulsifier was a soybean lecithin alone or in mixts. with Polysorbate 20. This nonionic surfactant was chosen since it was found to directly inhibit the hydrolysis of cefpodoxime proxetil in vitro using duodenal washings. The o/w submicron emulsions were be effective in protecting the prodrug from enzymic attack in rabbit duodenal washings compared with a micellar solution and an aqueous solution of cefpodoxime proxetil. An o/w submicron

emulsion incorporating Polysorbate 20 was e the most protective, which can corroborate the inhibitory role of Polysorbate itself.

IT 9005-64-5, Polysorbate 20 9005-70-3, Polysorbate 85

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(**cefpodoxime**-proxetil protection from intestinal lumen

hydrolysis by oil-in-water submicron emulsions)

REFERENCE COUNT: 41 THERE ARE 41 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L93 ANSWER 3 OF 3 HCAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER: 1983:204314 HCAPLUS

DOCUMENT NUMBER: 98:204314

TITLE: Effect of ethyl cellulose in a medium-chain triglyceride on the bioavailability of ceftizoxime

AUTHOR(S): Ueda, Ikuo; Shimojo, Fumio; Kozatani, Jun

CORPORATE SOURCE: Res. Lab., Fujisawa Pharm. Co., Ltd., Osaka, 532, Japan

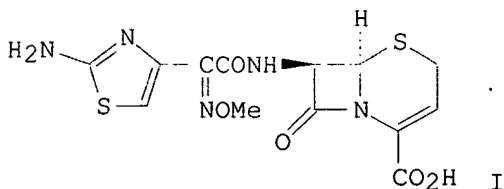
SOURCE: Journal of Pharmaceutical Sciences (1983), 72(4), 454-8

CODEN: JPMSAE; ISSN: 0022-3549

DOCUMENT TYPE: Journal

LANGUAGE: English

GI



AB The oral bioavailability of **ceftizoxime** Na (I Na) [ 68401-82-1] was improved by formulations containing Et cellulose [9004-57-3] and medium-chain triglycerides (Miglyol 812), as compared with aqueous solns. of I. A formulation containing 500 mg I Na, 5.0 mL medium-chain triglyceride, and 200 mg Et cellulose gave a larger area under the curve and higher urinary excretion of I than a formulation containing 50 mg Et cellulose instead of 200 mg. The urinary excretion of I was 7.1-10.3% when **polyethylene glycol**, olive oil, or H2O was used as the vehicle, as compared to 16.2-37.7% when the triglyceride + Et cellulose or olive oil + Et cellulose was used as the vehicle. Serum concns. attained exceeded the min. inhibitory concns. for most bacterial pathogens.

=> => sel hit rn 190 1-7  
E1 THROUGH E41 ASSIGNED

=> sel hit rn 191 1  
E42 THROUGH E42 ASSIGNED

=> sel hit rn 192 1-7  
E43 THROUGH E60 ASSIGNED

=> sel hit rn 193 1-3  
E61 THROUGH E63 ASSIGNED

=> file reg  
FILE 'REGISTRY' ENTERED AT 15:28:27 ON 03 FEB 2003  
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.  
PLEASE SEE "HELP USAGETERMS" FOR DETAILS.  
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STRUCTURE FILE UPDATES: 2 FEB 2003 HIGHEST RN 484639-64-7  
DICTIONARY FILE UPDATES: 2 FEB 2003 HIGHEST RN 484639-64-7

TSCA INFORMATION NOW CURRENT THROUGH MAY 20, 2002

Please note that search-term pricing does apply when  
conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. See HELP  
PROPERTIES for more information. See STNote 27, Searching Properties  
in the CAS Registry File, for complete details:  
<http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf>

=> => d his 194

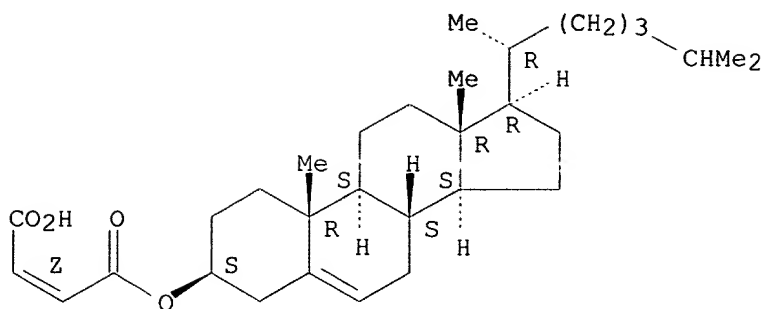
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SEL HIT RN L90 1-7  
SEL HIT RN L91 1  
SEL HIT RN L92 1-7  
SEL HIT RN L93 1-3

FILE 'REGISTRY' ENTERED AT 15:28:27 ON 03 FEB 2003  
L94 43 S E1-E63

=> d ide can 194 tot

L94 ANSWER 1 OF 43 REGISTRY COPYRIGHT 2003 ACS  
RN 133442-38-3 REGISTRY  
CN Cholest-5-en-3-ol (3 $\beta$ )-, hydrogen 2-butenedioate, lithium salt,  
[3(Z)]- (9CI) (CA INDEX NAME)  
FS STEREOSEARCH  
MF C31 H48 O4 . Li  
SR CA  
LC STN Files: CA, CAPLUS, TOXCENTER  
CRN (129357-50-2)

Absolute stereochemistry.  
Double bond geometry as shown.



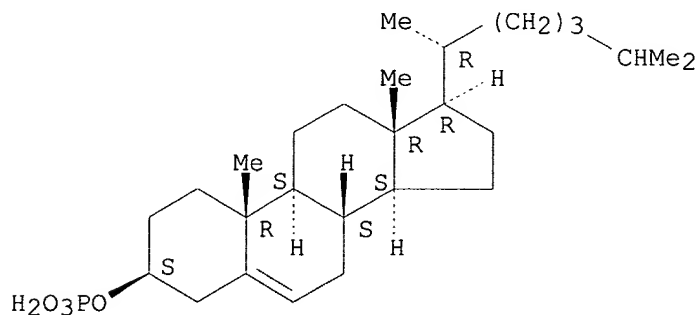
● Li

1 REFERENCES IN FILE CA (1962 TO DATE)  
1 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 114:192593

L94 ANSWER 2 OF 43 REGISTRY COPYRIGHT 2003 ACS  
RN 133352-86-0 REGISTRY  
CN Cholest-5-en-3-ol (3β)-, dihydrogen phosphate, calcium salt (9CI)  
(CA INDEX NAME)  
OTHER NAMES:  
CN Calcium cholesterol phosphate  
FS STEREOSEARCH  
MF C27 H47 O4 P . x Ca  
SR CA  
LC STN Files: CA, CAPLUS, TOXCENTER  
CRN (4358-16-1)

Absolute stereochemistry.



● x Ca

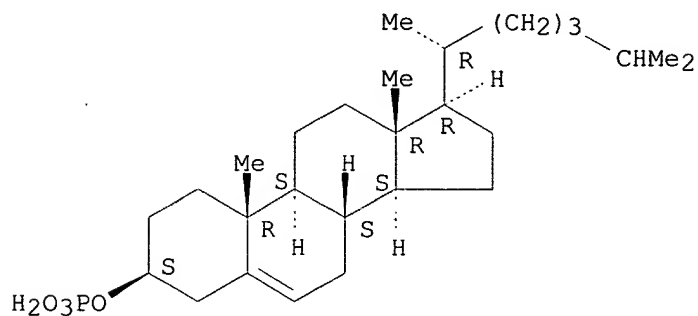
1 REFERENCES IN FILE CA (1962 TO DATE)  
1 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 114:192593

L94 ANSWER 3 OF 43 REGISTRY COPYRIGHT 2003 ACS  
RN 133352-85-9 REGISTRY  
CN Cholest-5-en-3-ol (3β)-, dihydrogen phosphate, magnesium salt (9CI)  
(CA INDEX NAME)  
OTHER NAMES:

CN Magnesium cholesterol phosphate  
 FS STEREOSEARCH  
 MF C27 H47 O4 P . x Mg  
 SR CA  
 LC STN Files: CA, CAPLUS, TOXCENTER  
 CRN (4358-16-1)

Absolute stereochemistry.



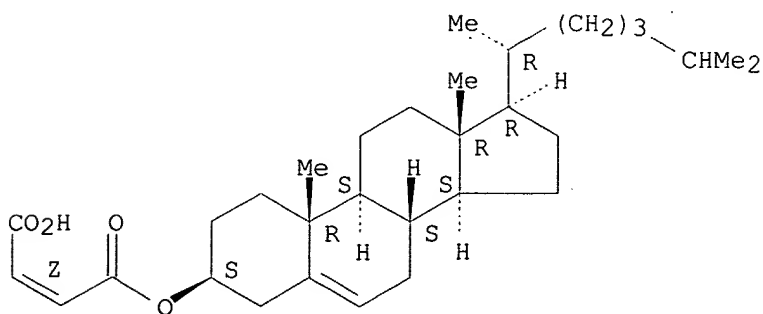
● x Mg

1 REFERENCES IN FILE CA (1962 TO DATE)  
 1 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 114:192593

L94 ANSWER 4 OF 43 REGISTRY COPYRIGHT 2003 ACS  
 RN 133161-26-9 REGISTRY  
 CN Cholest-5-en-3-ol (3β)-, hydrogen 2-butenedioate, calcium salt,  
 [3(Z)]- (9CI) (CA INDEX NAME)  
 FS STEREOSEARCH  
 MF C31 H48 O4 . 1/2 Ca  
 SR CA  
 LC STN Files: CA, CAPLUS, TOXCENTER  
 CRN (129357-50-2)

Absolute stereochemistry.  
 Double bond geometry as shown.



● 1/2 Ca

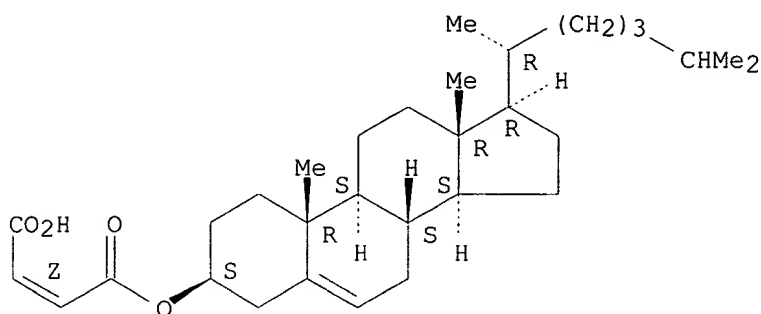
1 REFERENCES IN FILE CA (1962 TO DATE)  
 1 REFERENCES IN FILE CAPLUS (1962 TO DATE)



REFERENCE 1: 114:192593

L94 ANSWER 5 OF 43 REGISTRY COPYRIGHT 2003 ACS  
 RN 133161-25-8 REGISTRY  
 CN Cholest-5-en-3-ol (3 $\beta$ )-, hydrogen 2-butenedioate, magnesium salt,  
 [3(Z)]- (9CI) (CA INDEX NAME)  
 FS STEREOSEARCH  
 MF C31 H48 O4 . 1/2 Mg  
 SR CA  
 LC STN Files: CA, CAPLUS, TOXCENTER  
 CRN (129357-50-2)

Absolute stereochemistry.  
 Double bond geometry as shown.



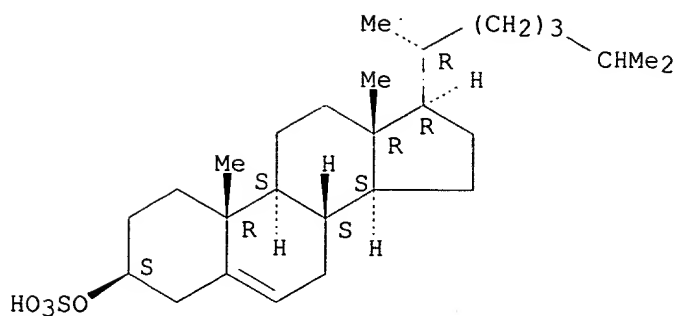
● 1/2 Mg

1 REFERENCES IN FILE CA (1962 TO DATE)  
 1 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 114:192593

L94 ANSWER 6 OF 43 REGISTRY COPYRIGHT 2003 ACS  
 RN 133058-05-6 REGISTRY  
 CN Cholest-5-en-3-ol (3 $\beta$ )-, hydrogen sulfate, magnesium salt (9CI) (CA  
 INDEX NAME)  
 OTHER NAMES:  
 CN Magnesium cholesteryl sulfate  
 FS STEREOSEARCH  
 MF C27 H46 O4 S . 1/2 Mg  
 SR CA  
 LC STN Files: CA, CAPLUS, TOXCENTER  
 CRN (1256-86-6)

Absolute stereochemistry.



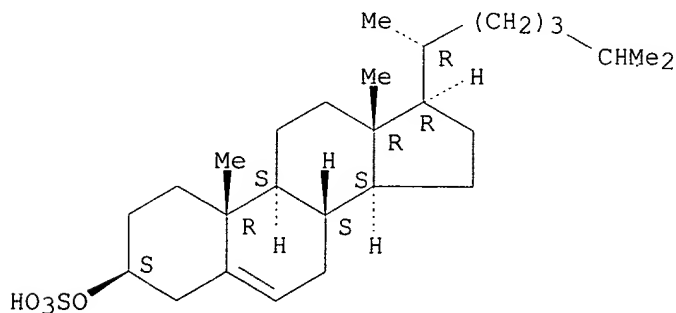
● 1/2 Mg

1 REFERENCES IN FILE CA (1962 TO DATE)  
1 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 114:192593

L94 ANSWER 7 OF 43 REGISTRY COPYRIGHT 2003 ACS  
RN 133058-04-5 REGISTRY  
CN Cholest-5-en-3-ol (3β)-, hydrogen sulfate, lithium salt (9CI) (CA INDEX NAME)  
FS STEREOSEARCH  
MF C27 H46 O4 S . Li  
SR CA  
LC STN Files: BEILSTEIN\*, CA, CAPLUS, TOXCENTER  
(\*File contains numerically searchable property data)  
CRN (1256-86-6)

Absolute stereochemistry.



● Li

1 REFERENCES IN FILE CA (1962 TO DATE)  
1 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 114:192593

L94 ANSWER 8 OF 43 REGISTRY COPYRIGHT 2003 ACS  
RN 113359-04-9 REGISTRY  
CN Imidazo[1,2-b]pyridazinium, 1-[[[(6R,7R)-7-[[[(2Z)-(5-amino-1,2,4-thiadiazol-3-yl)(methoxyimino)acetyl]amino]-2-carboxy-8-oxo-5-thia-1-azabicyclo[4.2.0]oct-2-en-3-yl]methyl]-, inner salt (9CI) (CA INDEX NAME)

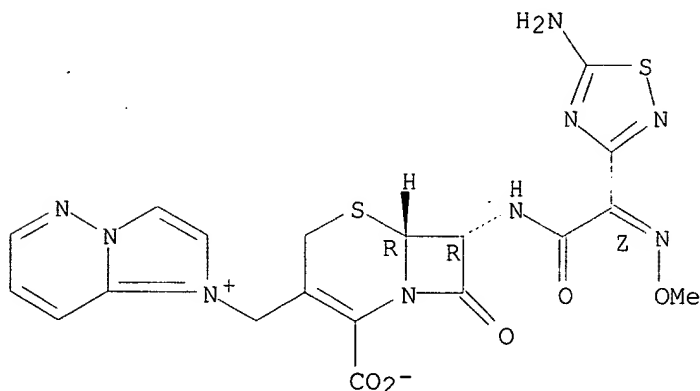
## OTHER CA INDEX NAMES:

CN 1,2,4-Thiadiazole, imidazo[1,2-b]pyridazinium deriv.  
 CN Imidazo[1,2-b]pyridazinium, 1-[[7-[[[(5-amino-1,2,4-thiadiazol-3-yl)(methoxyimino)acetyl]amino]-2-carboxy-8-oxo-5-thia-1-azabicyclo[4.2.0]oct-2-en-3-yl]methyl]-, inner salt, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]-

## OTHER NAMES:

CN Cefozopran  
 CN SCE 2787  
 FS STEREOSEARCH  
 DR 128007-70-5, 125882-76-0, 123572-82-7, 133790-04-2  
 MF C19 H17 N9 O5 S2  
 SR CA  
 LC STN Files: ADISINSIGHT, ADISNEWS, ANABSTR, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CANCERLIT, CAPLUS, CBNB, CIN, DDFU, DRUGNL, DRUGPAT, DRUGU, DRUGUPDATES, EMBASE, MEDLINE, MRCK\*, PHAR, PROMT, SYNTHLINE, TOXCENTER, USAN, USPATFULL  
 (\*File contains numerically searchable property data)  
 Other Sources: WHO

Absolute stereochemistry.  
 Double bond geometry as shown.



170 REFERENCES IN FILE CA (1962 TO DATE)

4 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

169 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 138:69776  
 REFERENCE 2: 138:69764  
 REFERENCE 3: 137:347477  
 REFERENCE 4: 137:291106  
 REFERENCE 5: 137:257189  
 REFERENCE 6: 137:244539  
 REFERENCE 7: 137:226242  
 REFERENCE 8: 137:152227  
 REFERENCE 9: 137:149790

REFERENCE 10: 137:114536

L94 ANSWER 9 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 105239-91-6 REGISTRY

CN 1-Azoniabicyclo[2.2.2]octane, 4-(aminocarbonyl)-1-[[[(6R,7R)-7-[[[(2Z)-(5-amino-1,2,4-thiadiazol-3-yl)(methoxyimino)acetyl]amino]-2-carboxy-8-oxo-5-thia-1-azabicyclo[4.2.0]oct-2-en-3-yl]methyl]-, inner salt (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 1,2,4-Thiadiazole, 1-azoniabicyclo[2.2.2]octane deriv.

CN 1-Azoniabicyclo[2.2.2]octane, 4-(aminocarbonyl)-1-[[7-[[[(5-amino-1,2,4-thiadiazol-3-yl)(methoxyimino)acetyl]amino]-2-carboxy-8-oxo-5-thia-1-azabicyclo[4.2.0]oct-2-en-3-yl]methyl]-, inner salt, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]-

OTHER NAMES:

CN Antibiotic E 1040

CN Cefclidin

CN Cefclidine

CN E 1040

CN E 1040 (antibiotic)

FS STEREOSEARCH

MF C21 H26 N8 O6 S2

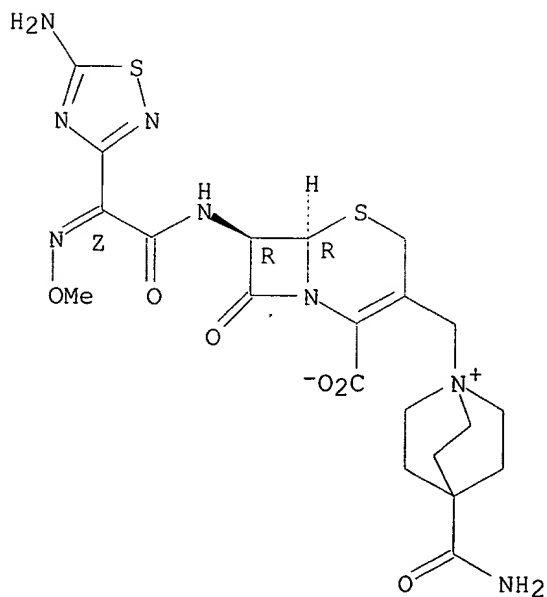
CI COM

SR CA

LC STN Files: ADISINSIGHT, BIOBUSINESS, BIOSIS, CA, CAPLUS, CIN, DDFU, DRUGNL, DRUGPAT, DRUGU, DRUGUPDATES, MRCK\*, PHAR, PROMT, RTECS\*, SYNTHLINE, TOXCENTER, USAN, USPATFULL  
(\*File contains numerically searchable property data)

Absolute stereochemistry.

Double bond geometry as shown.



102 REFERENCES IN FILE CA (1962 TO DATE)

4 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

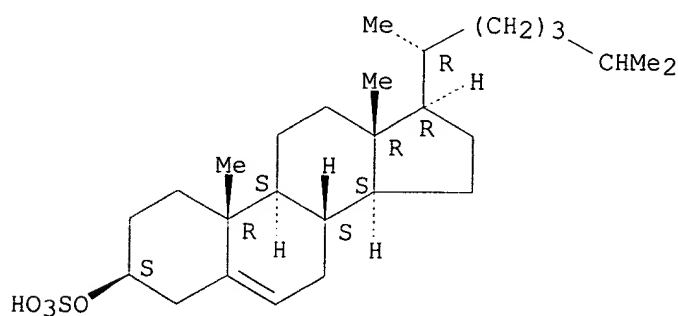
102 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 137:347477

REFERENCE 2: 137:341930  
 REFERENCE 3: 137:299611  
 REFERENCE 4: 137:149790  
 REFERENCE 5: 137:114536  
 REFERENCE 6: 136:172755  
 REFERENCE 7: 136:156464  
 REFERENCE 8: 136:156463  
 REFERENCE 9: 136:58849  
 REFERENCE 10: 134:222563

L94 ANSWER 10 OF 43 REGISTRY COPYRIGHT 2003 ACS  
 RN 99523-97-4 REGISTRY  
 CN Cholest-5-en-3-ol (3 $\beta$ )-, hydrogen sulfate, calcium salt (9CI) (CA INDEX NAME)  
 OTHER NAMES:  
 CN Calcium cholesteryl sulfate  
 CN Cholesterol hydrogen sulfate calcium salt  
 CN Cholesteryl sulfate calcium salt  
 FS STEREOSEARCH  
 DR 127565-96-2  
 MF C27 H46 O4 S . 1/2 Ca  
 SR CA  
 LC STN Files: BEILSTEIN\*, CA, CAPLUS, TOXCENTER  
 (\*File contains numerically searchable property data)  
 CRN (1256-86-6)

Absolute stereochemistry.



● 1/2 Ca

5 REFERENCES IN FILE CA (1962 TO DATE)  
 5 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 115:166374  
 REFERENCE 2: 114:192593  
 REFERENCE 3: 113:11949

REFERENCE 4: 113:11932

REFERENCE 5: 104:10400

L94 ANSWER 11 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 84957-29-9 REGISTRY

CN 5H-Cyclopenta[b]pyridinium, 1-[[[(6R,7R)-7-[[[(2Z)-(2-amino-4-thiazolyl)(methoxyimino)acetyl]amino]-2-carboxy-8-oxo-5-thia-1-azabicyclo[4.2.0]oct-2-en-3-yl]methyl]-6,7-dihydro-, inner salt (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 5H-1-Pyridinium, 1-[[[7-[[[(2-amino-4-thiazolyl)(methoxyimino)acetyl]amino]-2-carboxy-8-oxo-5-thia-1-azabicyclo[4.2.0]oct-2-en-3-yl]methyl]-6,7-dihydro-, inner salt, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]-

OTHER NAMES:

CN 5H-Cyclopenta[b]pyridinium, 1-[[[7-[[[(2-amino-4-thiazolyl)(methoxyimino)acetyl]amino]-2-carboxy-8-oxo-5-thia-1-azabicyclo[4.2.0]oct-2-en-3-yl]methyl]-6,7-dihydro-, inner salt, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]-

CN Antibiotic HR 810

CN Cefpirome

CN Cefrom

CN HR 810

FS STEREOSEARCH

MF C22 H22 N6 O5 S2

CI COM

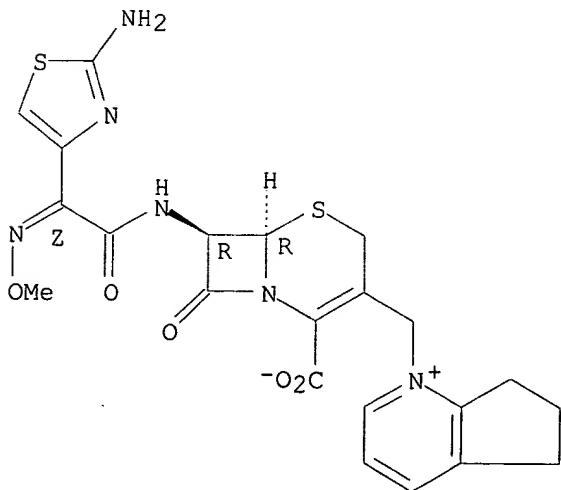
LC STN Files: ADISINSIGHT, ADISNEWS, ANABSTR, BEILSTEIN\*, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CANCERLIT, CAPLUS, CASREACT, CBNB, CIN, DDFU, DRUGNL, DRUGPAT, DRUGU, DRUGUPDATES, EMBASE, IPA, MEDLINE, MRCK\*, PHAR, PROMT, SYNTHLINE, TOXCENTER, USAN, USPATFULL

(\*File contains numerically searchable property data)

Other Sources: WHO

Absolute stereochemistry.

Double bond geometry as shown.



591 REFERENCES IN FILE CA (1962 TO DATE)

6 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

593 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 138:69776

REFERENCE 2: 138:69764

REFERENCE 3: 138:51770  
 REFERENCE 4: 138:2166  
 REFERENCE 5: 137:347477  
 REFERENCE 6: 137:259865  
 REFERENCE 7: 137:163290  
 REFERENCE 8: 137:152227  
 REFERENCE 9: 137:149790  
 REFERENCE 10: 137:121424

L94 ANSWER 12 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 80370-57-6 REGISTRY

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
 7-[[[(2Z)-(2-amino-4-thiazolyl)(methoxyimino)acetyl]amino]-3-[[[(2-  
 furanylcarbonyl)thio]methyl]-8-oxo-, (6R,7R)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
 7-[[[(2-amino-4-thiazolyl)(methoxyimino)acetyl]amino]-3-[[[(2-  
 furanylcarbonyl)thio]methyl]-8-oxo-, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]-

OTHER NAMES:

CN Ceftiofur

CN CM 31916

FS STEREOSEARCH

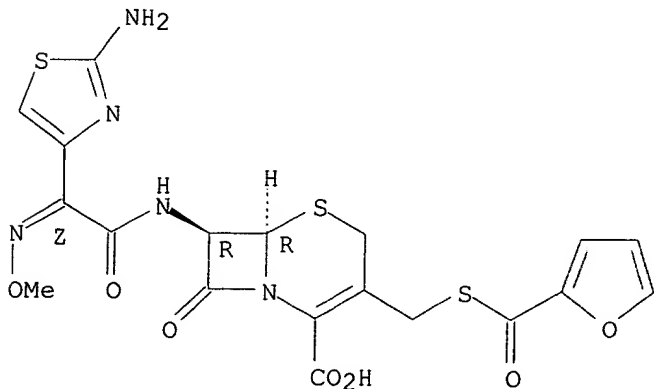
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CI COM

LC STN Files: AGRICOLA, ANABSTR, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CABA,  
 CANCERLIT, CAPLUS, CASREACT, CBNB, CIN, DDFU, DIOGENES, DRUGU, EMBASE,  
 MEDLINE, MRCK\*, PROMT, TOXCENTER, USAN, USPAT2, USPATFULL, VETU  
 (\*File contains numerically searchable property data)

Other Sources: WHO

Absolute stereochemistry.  
 Double bond geometry as shown.



\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

191 REFERENCES IN FILE CA (1962 TO DATE)

7 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
192 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 138:69785  
REFERENCE 2: 138:32649  
REFERENCE 3: 137:384233  
REFERENCE 4: 137:322471  
REFERENCE 5: 137:309620  
REFERENCE 6: 137:241635  
REFERENCE 7: 137:226224  
REFERENCE 8: 137:200279  
REFERENCE 9: 137:198165  
REFERENCE 10: 137:190750

L94 ANSWER 13 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 80210-62-4 REGISTRY

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
7-[[ (2Z)-(2-amino-4-thiazolyl) (methoxyimino)acetyl]amino]-3-  
(methoxymethyl)-8-oxo-, (6R,7R)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
7-[[ (2-amino-4-thiazolyl) (methoxyimino)acetyl]amino]-3-(methoxymethyl)-8-  
oxo-, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]-

OTHER NAMES:

CN Cefpodoxime

CN Cefpodoxime acid

CN R 3763

FS STEREOSEARCH

MF C15 H17 N5 O6 S2

CI COM

LC STN Files: ADISNEWS, ANABSTR, BEILSTEIN\*, BIOBUSINESS, BIOSIS, CA,  
CAPLUS, CASREACT, CBNB, CHEMINFORMRX, CIN, DDFU, DIOGENES, DRUGPAT,  
DRUGU, DRUGUPDATES, EMBASE, IPA, MEDLINE, PHAR, PROMT, TOXCENTER, USAN,  
USPAT2, USPATFULL

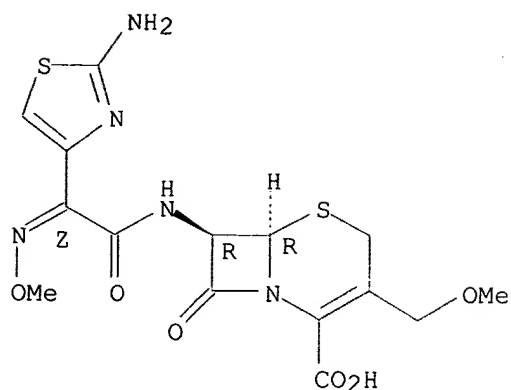
(\*File contains numerically searchable property data)

Other Sources: WHO

Absolute stereochemistry.

Double bond geometry as shown.





\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

318 REFERENCES IN FILE CA (1962 TO DATE)  
 2 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
 318 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 138:69776  
 REFERENCE 2: 138:52242  
 REFERENCE 3: 138:44695  
 REFERENCE 4: 138:22020  
 REFERENCE 5: 137:382147  
 REFERENCE 6: 137:366241  
 REFERENCE 7: 137:349153  
 REFERENCE 8: 137:349152  
 REFERENCE 9: 137:335069  
 REFERENCE 10: 137:322531

L94 ANSWER 14 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 79350-37-1 REGISTRY

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
 7-[[ (2Z)-(2-amino-4-thiazolyl) [(carboxymethoxy)imino]acetyl]amino]-3-  
 ethenyl-8-oxo-, (6R,7R)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
 7-[[ (2-amino-4-thiazolyl) [(carboxymethoxy)imino]acetyl]amino]-3-ethenyl-8-  
 oxo-, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]-

OTHER NAMES:

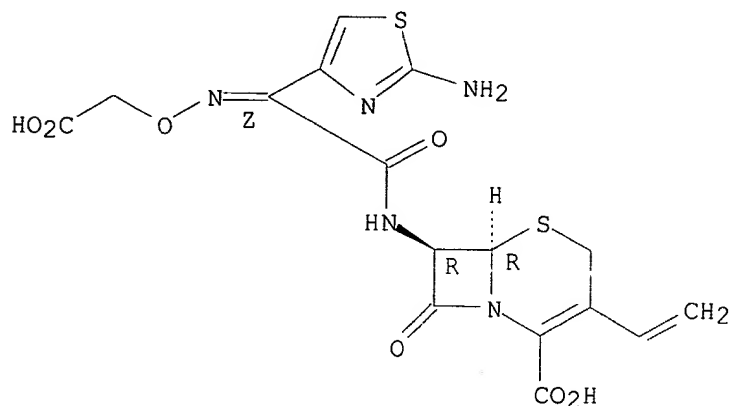
CN Cefixime  
 CN Cephoral  
 CN CL 284635  
 CN FK 027  
 CN FR 17027  
 CN Oroken  
 CN Suprax  
 FS STEREOSEARCH  
 DR 214265-67-5

MF C16 H15 N5 O7 S2

CI COM

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, BEILSTEIN\*, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CANCERLIT, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS, CIN, CSCHEM, DDFU, DIOGENES, DRUGNL, DRUGPAT, DRUGU, DRUGUPDATES, EMBASE, IPA, MEDLINE, MRCK\*, MSDS-OHS, PHAR, PHARMASEARCH, PROMT, RTECS\*, SYNTHLINE, TOXCENTER, USAN, USPAT2, USPATFULL, VETU  
 (\*File contains numerically searchable property data)  
 Other Sources: WHO

Absolute stereochemistry.  
 Double bond geometry as shown.



\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

717 REFERENCES IN FILE CA (1962 TO DATE)  
 7 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
 717 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 138:52587  
 REFERENCE 2: 138:29161  
 REFERENCE 3: 138:19297  
 REFERENCE 4: 137:382158  
 REFERENCE 5: 137:366241  
 REFERENCE 6: 137:349153  
 REFERENCE 7: 137:349152  
 REFERENCE 8: 137:347477  
 REFERENCE 9: 137:346142  
 REFERENCE 10: 137:335069

L94 ANSWER 15 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 74578-69-1 REGISTRY

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
 7-[[[(2Z)-(2-amino-4-thiazolyl)(methoxyimino)acetyl]amino]-8-oxo-3-  
 [[[(1,2,5,6-tetrahydro-2-methyl-5,6-dioxo-1,2,4-triazin-3-yl)thio]methyl]]-,  
 disodium salt, (6R,7R)- (9CI) (CA INDEX NAME)

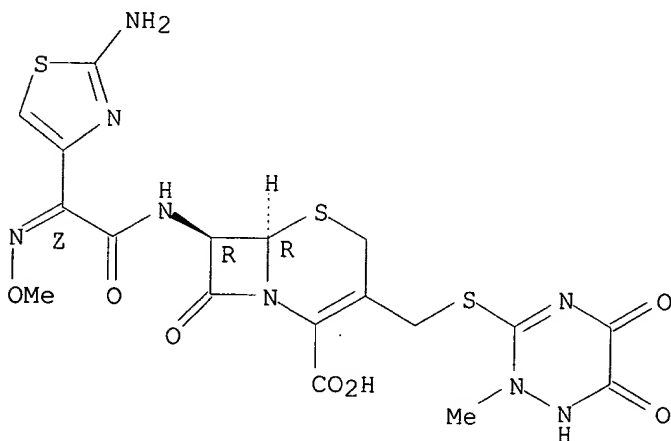
OTHER CA INDEX NAMES:

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
7-[[[(2-amino-4-thiazolyl)(methoxyimino)acetyl]amino]-8-oxo-3-[[[(1,2,5,6-tetrahydro-2-methyl-5,6-dioxo-1,2,4-triazin-3-yl)thio]methyl]-, disodium  
salt, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]-

## OTHER NAMES:

CN Ceftriaxone  
CN Ceftriaxone disodium  
CN Ceftriaxone sodium  
CN Cephtriaxone  
CN Longaceph  
CN Ro 13-9904  
CN Rocephin  
CN X 13-9904  
FS STEREOSEARCH  
MF C18 H18 N8 O7 S3 . 2 Na  
CI COM  
LC STN Files: ADISNEWS, ANABSTR, BEILSTEIN\*, BIOBUSINESS, BIOSIS,  
BIOTECHNO, CA, CABA, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS, CHEMLIST,  
CIN, CSCHEM, DIOGENES, DRUGPAT, DRUGUPDATES, EMBASE, IPA, MSDS-OHS,  
PHAR, PHARMASEARCH, PROMT, RTECS\*, SYNTHLINE, TOXCENTER, USPAT2,  
USPATFULL  
(\*File contains numerically searchable property data)  
Other Sources: EINECS\*\*  
(\*\*Enter CHEMLIST File for up-to-date regulatory information)  
CRN (73384-59-5)

Absolute stereochemistry.  
Double bond geometry as shown.



● 2 Na

183 REFERENCES IN FILE CA (1962 TO DATE)  
2 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
183 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 137:362547  
REFERENCE 2: 137:257592  
REFERENCE 3: 137:154929  
REFERENCE 4: 137:119619

REFERENCE 5: 137:711  
 REFERENCE 6: 136:369552  
 REFERENCE 7: 136:355103  
 REFERENCE 8: 136:319352  
 REFERENCE 9: 136:304035  
 REFERENCE 10: 136:90988

L94 ANSWER 16 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 73547-61-2 REGISTRY

CN Pyridinium, 1-[[[(6R,7R)-7-[[[(2Z)-(2-amino-4-thiazolyl)][(1-carboxy-1-methylethoxy)imino]acetyl]amino]-2-carboxy-8-oxo-5-thia-1-azabicyclo[4.2.0]oct-2-en-3-yl]methyl]-, inner salt, monosodium salt (9CI)  
 (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Pyridinium, 1-[[[7-[[[(2-amino-4-thiazolyl)][(1-carboxy-1-methylethoxy)imino]acetyl]amino]-2-carboxy-8-oxo-5-thia-1-azabicyclo[4.2.0]oct-2-en-3-yl]methyl]-, inner salt, monosodium salt, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]-

OTHER NAMES:

CN Ceftazidime sodium

CN GR 20263 monosodium salt

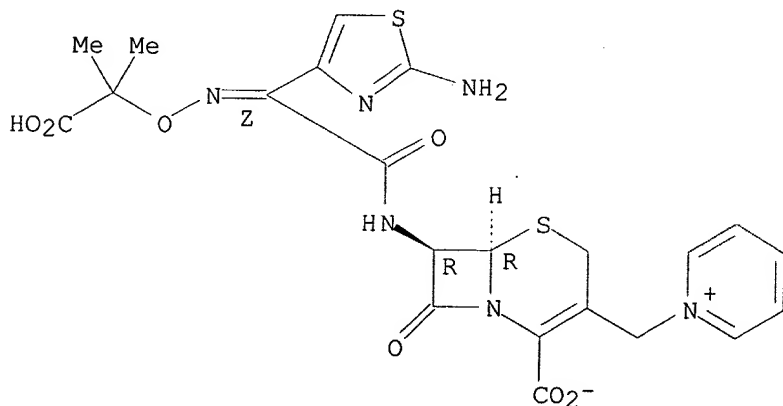
FS STEREOSEARCH

MF C22 H22 N6 O7 S2 . Na

LC STN Files: CA, CAPLUS, CIN, DIOGENES, DRUGPAT, IPA, MSDS-OHS, TOXCENTER, USPATFULL

CRN (72558-82-8)

Absolute stereochemistry.  
 Double bond geometry as shown.



● Na

15 REFERENCES IN FILE CA (1962 TO DATE)  
 15 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 135:170587

REFERENCE 2: 130:335196  
REFERENCE 3: 128:110395  
REFERENCE 4: 127:214608  
REFERENCE 5: 127:90147  
REFERENCE 6: 126:135559  
REFERENCE 7: 126:36948  
REFERENCE 8: 112:104629  
REFERENCE 9: 105:120614  
REFERENCE 10: 101:78747

L94 ANSWER 17 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 73384-59-5 REGISTRY

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
7-[[[(2Z)-(2-amino-4-thiazolyl)(methoxyimino)acetyl]amino]-8-oxo-3-  
[[[(1,2,5,6-tetrahydro-2-methyl-5,6-dioxo-1,2,4-triazin-3-yl)thio]methyl]-,  
(6R,7R)-(9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
7-[[[(2-amino-4-thiazolyl)(methoxyimino)acetyl]amino]-8-oxo-3-[[[(1,2,5,6-  
tetrahydro-2-methyl-5,6-dioxo-1,2,4-triazin-3-yl)thio]methyl]-,  
[6R-[6 $\alpha$ ,7 $\beta$ (Z)]]-

OTHER NAMES:

CN Biotrakson

CN Ceftriaxone

CN Rocefin

FS STEREOSEARCH

DR 380149-23-5

MF C18 H18 N8 O7 S3

CI COM

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, BEILSTEIN\*, BIOBUSINESS, BIOSIS,  
BIOTECHNO, CA, CABA, CANCERLIT, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS,  
CHEMLIST, CIN, CSCHEM, CSNB, DDFU, DRUGNL, DRUGPAT, DRUGU, DRUGUPDATES,  
EMBASE, IFICDB, IFIUDB, IPA, MEDLINE, MRCK\*, NIOSHTIC, PHAR,  
PHARMASEARCH, PROMT, RTECS\*, SYNTHLINE, TOXCENTER, USAN, USPAT2,  
USPATFULL, VETU

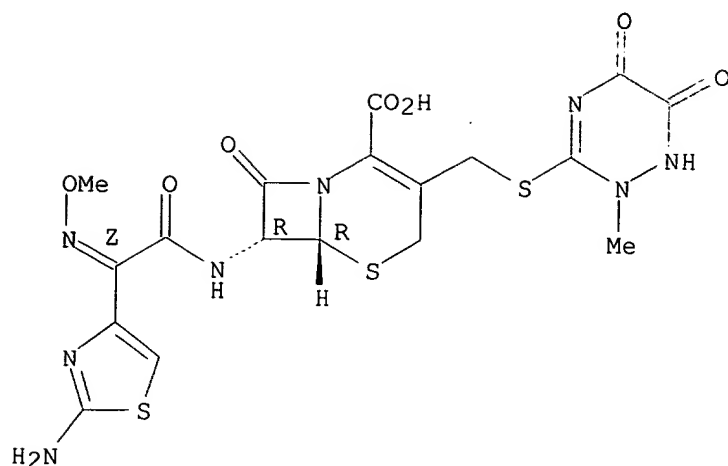
(\*File contains numerically searchable property data)

Other Sources: EINECS\*\*, WHO

(\*\*Enter CHEMLIST File for up-to-date regulatory information)

Absolute stereochemistry.

Double bond geometry as shown.



\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

2376 REFERENCES IN FILE CA (1962 TO DATE)  
 15 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
 2382 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 138:78450

REFERENCE 2: 138:70922

REFERENCE 3: 138:69793

REFERENCE 4: 138:69790

REFERENCE 5: 138:69776

REFERENCE 6: 138:66192

REFERENCE 7: 138:66102

REFERENCE 8: 138:52614

REFERENCE 9: 138:52597

REFERENCE 10: 138:52587

L94 ANSWER 18 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 72558-82-8 REGISTRY

CN Pyridinium, 1-[[[(6R, 7R)-7-[[[(2Z)-(2-amino-4-thiazolyl)][(1-carboxy-1-methylethoxy)imino]acetyl]amino]-2-carboxy-8-oxo-5-thia-1-azabicyclo[4.2.0]oct-2-en-3-yl)methyl]-, inner salt (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Pyridinium, 1-[[7-[[[(2-amino-4-thiazolyl)][(1-carboxy-1-methylethoxy)imino]acetyl]amino]-2-carboxy-8-oxo-5-thia-1-azabicyclo[4.2.0]oct-2-en-3-yl)methyl]-, hydroxide, inner salt, [6R-[6 $\alpha$ , 7 $\beta$ (Z)]]-

OTHER NAMES:

CN (6R, 7R)-7-[(Z)-2-(Aminothiazol-4-yl)-2-(2-carboxypropoxyimino)acetamido]-3-(1-pyridiniummethyl)ceph-3-em-4-carboxylate

CN Biotum

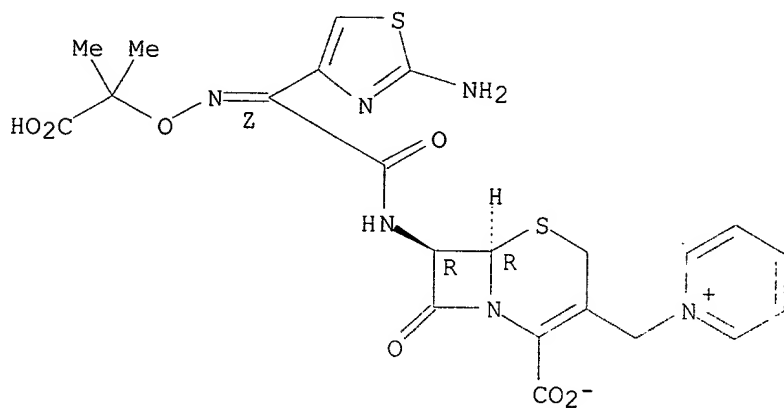
CN Ceftacidin

CN Ceftazidime

CN Ceftazidine  
 CN Fortaz  
 CN Fortum  
 CN GR 20263  
 CN Modacin  
 CN Tazicef  
 CN Tazidime  
 FS STEREOSEARCH  
 DR 133943-58-5  
 MF C22 H22 N6 O7 S2  
 CI COM

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, BEILSTEIN\*, BIOBUSINESS, BIOSIS,  
 BIOTECHNO, CA, CABA, CAPLUS, CASREACT, CBNB, CEN, CHEMLIST, CIN, CSCHM,  
 CSNB, DDFU, DIOGENES, DRUGPAT, DRUGU, EMBASE, IFICDB, IFIUDB, MRCK\*,  
 MSDS-OHS, NIOSHTIC, PHAR, PHARMASEARCH, PIRA, PROMT, RTECS\*, TOXCENTER,  
 USPAT2, USPATFULL, VETU  
 (\*File contains numerically searchable property data)  
 Other Sources: EINECS\*\*, WHO  
 (\*\*Enter CHEMLIST File for up-to-date regulatory information)

Absolute stereochemistry.  
 Double bond geometry as shown.



3419 REFERENCES IN FILE CA (1962 TO DATE)  
 16 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
 3426 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 138:69776  
 REFERENCE 2: 138:69764  
 REFERENCE 3: 138:69353  
 REFERENCE 4: 138:69078  
 REFERENCE 5: 138:66192  
 REFERENCE 6: 138:65570  
 REFERENCE 7: 138:61354  
 REFERENCE 8: 138:52593  
 REFERENCE 9: 138:51836  
 REFERENCE 10: 138:51770

L94 ANSWER 19 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 68401-82-1 REGISTRY

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
7-[[[(2Z)-(2-amino-4-thiazolyl)(methoxyimino)acetyl]amino]-8-oxo-,  
monosodium salt, (6R,7R)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
7-[[[(2-amino-4-thiazolyl)(methoxyimino)acetyl]amino]-8-oxo-, monosodium  
salt, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]-

OTHER NAMES:

CN Ceftizoxime sodium

CN Ceftizoxime sodium salt

CN FK 749

CN Sodium 7-[2-(2-aminothiazol-4-yl)-2-methoxyiminoacetamido]-3-cephem-4-  
carboxylate

FS STEREOSEARCH

MF C13 H13 N5 O5 S2 . Na

CI COM

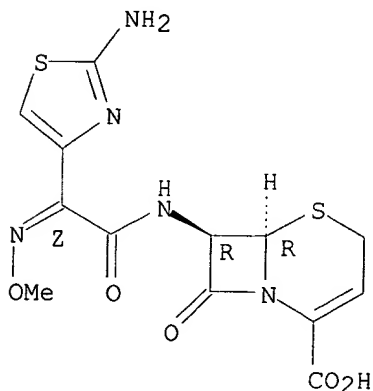
LC STN Files: BEILSTEIN\*, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CAPLUS,  
CASREACT, CBNB, CHEMLIST, CIN, DIOGENES, DRUGPAT, EMBASE, IFICDB,  
IFIUDB, IPA, MRCK\*, MSDS-OHS, PHAR, PROMT, RTECS\*, TOXCENTER, USAN,  
USPATFULL

(\*File contains numerically searchable property data)

CRN (68401-81-0)

Absolute stereochemistry.

Double bond geometry as shown.



● Na

88 REFERENCES IN FILE CA (1962 TO DATE)

1 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

88 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 137:217244

REFERENCE 2: 136:11106

REFERENCE 3: 135:170587

REFERENCE 4: 135:51028

REFERENCE 5: 135:37082



REFERENCE 6: 134:290352

REFERENCE 7: 134:242521

REFERENCE 8: 131:170633

REFERENCE 9: 128:110395

REFERENCE 10: 127:325983

L94 ANSWER 20 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 68401-81-0 REGISTRY

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
7-[[ (2Z)-(2-amino-4-thiazolyl) (methoxyimino)acetyl]amino]-8-oxo-, (6R,7R)-  
(9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
7-[[ (2-amino-4-thiazolyl) (methoxyimino)acetyl]amino]-8-oxo-,  
[6R-[6 $\alpha$ ,7 $\beta$ (Z)]]-

OTHER NAMES:

CN Ceftisomin

CN Ceftizoxime

CN Epocelin

FS STEREOSEARCH

MF C13 H13 N5 O5 S2

CI COM

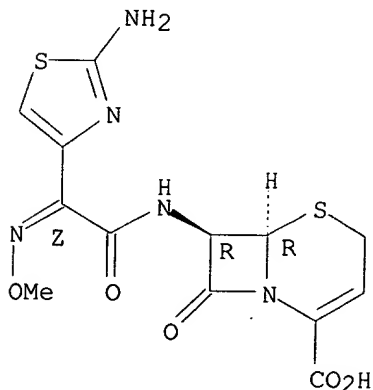
LC STN Files: ADISINSIGHT, ADISNEWS, AGRICOLA, ANABSTR, BEILSTEIN\*,  
BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CANCERLIT, CAPLUS, CASREACT, CBNB,  
CIN, DDFU, DIOGENES, DRUGPAT, DRUGU, EMBASE, IFICDB, IFIUDB, IPA,  
MEDLINE, MRCK\*, NIOSHTIC, PHAR, PROMT, RTECS\*, TOXCENTER, USAN, USPAT2,  
USPATFULL, VETU

(\*File contains numerically searchable property data)

Other Sources: WHO

Absolute stereochemistry.

Double bond geometry as shown.



\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

979 REFERENCES IN FILE CA (1962 TO DATE)

4 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

979 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 138:69780  
REFERENCE 2: 138:49428  
REFERENCE 3: 138:12452  
REFERENCE 4: 137:347477  
REFERENCE 5: 137:237750  
REFERENCE 6: 137:226224  
REFERENCE 7: 137:210914  
REFERENCE 8: 137:169369  
REFERENCE 9: 137:166077  
REFERENCE 10: 137:149790

L94 ANSWER 21 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 65085-01-0 REGISTRY

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
7-[[[(2Z)-(2-amino-4-thiazolyl)(methoxyimino)acetyl]amino]-3-[[[(1-methyl-1H-  
tetrazol-5-yl)thio]methyl]-8-oxo-, (6R,7R)-(9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
7-[[[(2-amino-4-thiazolyl)(methoxyimino)acetyl]amino]-3-[[[(1-methyl-1H-  
tetrazol-5-yl)thio]methyl]-8-oxo-, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]-

OTHER NAMES:

CN AB 50912

CN Cefmenoxime

CN SCE 1365

FS STEREOSEARCH

DR 74512-79-1, 74565-79-0

MF C16 H17 N9 O5 S3

CI COM

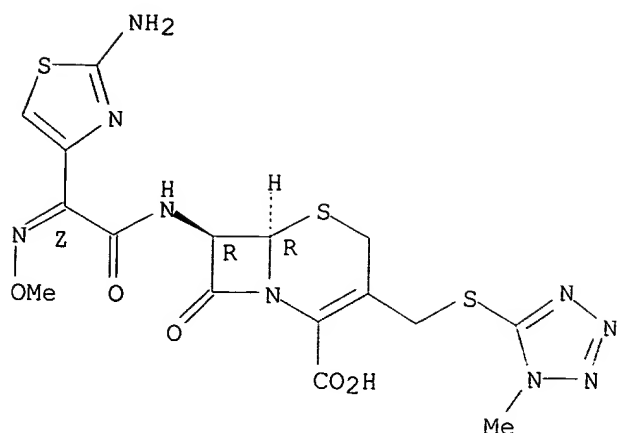
LC STN Files: ADISNEWS, ANABSTR, BEILSTEIN\*, BIOBUSINESS, BIOSIS,  
BIOTECHNO, CA, CANCERLIT, CAPLUS, CASREACT, CBNB, CIN, DDFU, DRUGPAT,  
DRUGU, EMBASE, IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, MRCK\*, PHAR, PROMT,  
TOXCENTER, USAN, USPATFULL

(\*File contains numerically searchable property data)

Other Sources: WHO

Absolute stereochemistry.

Double bond geometry as shown.



\*\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*\*

538 REFERENCES IN FILE CA (1962 TO DATE)  
7 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
538 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 137:347477

REFERENCE 2: 137:149790

REFERENCE 3: 137:114536

REFERENCE 4: 137:90836

REFERENCE 5: 136:172755

REFERENCE 6: 136:156464

REFERENCE 7: 136:156463

REFERENCE 8: 136:123678

REFERENCE 9: 136:58849

REFERENCE 10: 135:73913

L94 ANSWER 22 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 63527-52-6 REGISTRY

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
3-[(acetyloxy)methyl]-7-[[ (2Z)-(2-amino-4-thiazolyl) (methoxyimino)acetyl]a  
mino]-8-oxo-, (6R,7R)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
3-[(acetyloxy)methyl]-7-[[[(2-amino-4-thiazolyl)(methoxyimino)acetyl]amino]-  
8-oxo-, [6R-[6 $\alpha$ ,7 $\beta$ (Z)]]-

OTHER NAMES:

CN Cefotaxime

CN Cephotaxime

CN Claforan

FS STEREOSEARCH

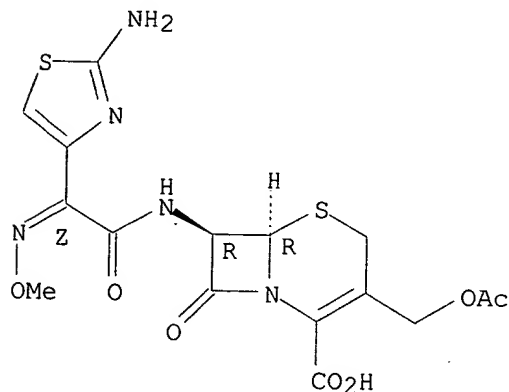
MF C16 H17 N5 O7 S2

CI	COM
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LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, BEILSTEIN\*, BIOBUSINESS, BIOSIS,  
BIOTECHNO, CA, CANCERLIT, CAPLUS, CASREACT, CBNB, CEN, CHEMLIST, CIN,

CSCHEM, DDFU, DIOGENES, DRUGPAT, DRUGU, EMBASE, IFICDB, IFIPAT, IFIUDB,  
IPA, MEDLINE, MRCK\*, NAPRALERT, NIOSHTIC, PHARMASEARCH, PROMT, RTECS\*,  
SYNTHLINE, TOXCENTER, USAN, USPAT2, USPATFULL, VETU  
(\*File contains numerically searchable property data)  
Other Sources: EINECS\*\*, WHO  
(\*\*Enter CHEMLIST File for up-to-date regulatory information)

Absolute stereochemistry.  
Double bond geometry as shown.



\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

4548 REFERENCES IN FILE CA (1962 TO DATE)  
32 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
4555 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 138:69776  
REFERENCE 2: 138:69767  
REFERENCE 3: 138:69078  
REFERENCE 4: 138:66192  
REFERENCE 5: 138:65570  
REFERENCE 6: 138:52593  
REFERENCE 7: 138:52588  
REFERENCE 8: 138:52242  
REFERENCE 9: 138:51770  
REFERENCE 10: 138:35956

L94 ANSWER 23 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 62893-20-3 REGISTRY

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
7-[[[(2R)-[[[(4-ethyl-2,3-dioxo-1-piperazinyl)carbonyl]amino](4-  
hydroxyphenyl)acetyl]amino]-3-[[[(1-methyl-1H-tetrazol-5-yl)thio]methyl]-8-  
oxo-, monosodium salt, (6R,7R)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,

7-[[[(4-ethyl-2,3-dioxo-1-piperazinyl)carbonyl]amino](4-hydroxyphenyl)acetyl]amino]-3-[[[(1-methyl-1H-tetrazol-5-yl)thio]methyl]-8-oxo-, monosodium salt, [6R-[6 $\alpha$ ,7 $\beta$ (R\*)]]-

## OTHER NAMES:

CN Cefoperazone sodium

CN Cefoperazone sodium salt

CN Sodium cefoperazone

CN T 1551

FS STEREOSEARCH

MF C25 H27 N9 O8 S2 . Na

CI COM

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, BEILSTEIN\*, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CABA, CAPLUS, CASREACT, CBNB, CHEMCATS, CHEMLIST, CIN, CSChem, DIOGENES, DRUGPAT, EMBASE, IPA, MRCK\*, PROMT, RTECS\*, SYNTHLINE, TOXCENTER, USAN, USPATFULL

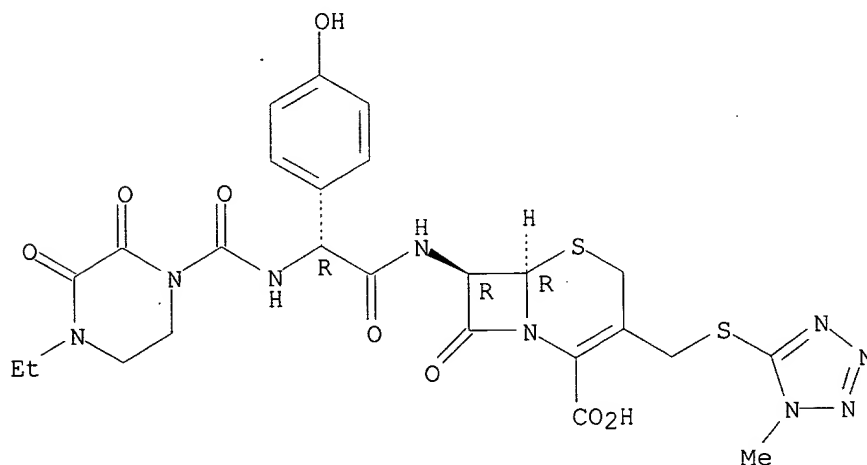
(\*File contains numerically searchable property data)

Other Sources: EINECS\*\*

(\*\*Enter CHEMLIST File for up-to-date regulatory information)

CRN (62893-19-0)

Absolute stereochemistry.



● Na

111 REFERENCES IN FILE CA (1962 TO DATE)

1 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

111 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 137:362547

REFERENCE 2: 137:329562

REFERENCE 3: 137:175107

REFERENCE 4: 137:145696

REFERENCE 5: 136:304035

REFERENCE 6: 135:318356

REFERENCE 7: 135:51028

REFERENCE 8: 135:37082

REFERENCE 9: 134:27272

REFERENCE 10: 133:237755

L94 ANSWER 24 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 62893-19-0 REGISTRY

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
 7-[[[(2R)-[[[(4-ethyl-2,3-dioxo-1-piperazinyl)carbonyl]amino](4-  
 hydroxyphenyl)acetyl]amino]-3-[[[(1-methyl-1H-tetrazol-5-yl)thio]methyl]-8-  
 oxo-, (6R,7R)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 5-Thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid,  
 7-[[[[[(4-ethyl-2,3-dioxo-1-piperazinyl)carbonyl]amino](4-  
 hydroxyphenyl)acetyl]amino]-3-[[[(1-methyl-1H-tetrazol-5-yl)thio]methyl]-8-  
 oxo-, [6R-[6 $\alpha$ ,7 $\beta$ (R\*)]]-

OTHER NAMES:

CN Cefob

CN Cefobid

CN Cefoperazine

CN Cefoperazone

CN Cephaperazon

CN Medocef

FS STEREOSEARCH

DR 72448-63-6

MF C25 H27 N9 O8 S2

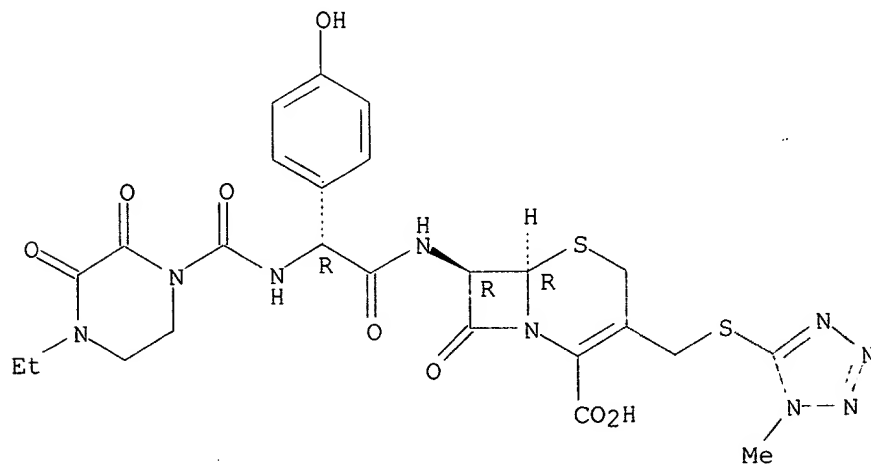
CI COM

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, BEILSTEIN\*, BIOBUSINESS, BIOSIS,  
 BIOTECHNO, CA, CABA, CANCERLIT, CAPLUS, CASREACT, CBNB, CHEMCATS,  
 CHEMLIST, CIN, CSCHEM, DDFU, DIOGENES, DRUGPAT, DRUGU, EMBASE, IFICDB,  
 IFIUDB, IPA, MEDLINE, MRCK\*, PHAR, PHARMASEARCH, PROMT, RTECS\*,  
 SYNTHLINE, TOXCENTER, USAN, USPAT2, USPATFULL, VETU  
 (\*File contains numerically searchable property data)

Other Sources: EINECS\*\*, WHO

(\*\*Enter CHEMLIST File for up-to-date regulatory information)

Absolute stereochemistry.



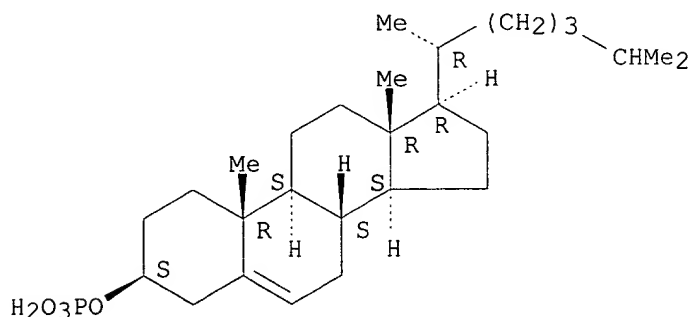
\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

2038 REFERENCES IN FILE CA (1962 TO DATE)  
 13 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
 2042 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 138:69776  
 REFERENCE 2: 138:66192  
 REFERENCE 3: 138:44717  
 REFERENCE 4: 138:11124  
 REFERENCE 5: 138:2166  
 REFERENCE 6: 137:382143  
 REFERENCE 7: 137:366212  
 REFERENCE 8: 137:351600  
 REFERENCE 9: 137:347477  
 REFERENCE 10: 137:336792

L94 ANSWER 25 OF 43 REGISTRY COPYRIGHT 2003 ACS  
 RN 24352-55-4 REGISTRY  
 CN Cholest-5-en-3-ol (3 $\beta$ )-, dihydrogen phosphate, dilithium salt (9CI)  
 (CA INDEX NAME)  
 OTHER CA INDEX NAMES:  
 CN Cholesterol, dihydrogen phosphate, dilithium salt (8CI)  
 FS STEREOSEARCH  
 MF C27 H47 O4 P . 2 Li  
 LC STN Files: CA, CAPLUS, TOXCENTER  
 CRN (4358-16-1)

Absolute stereochemistry.



● 2 Li

2 REFERENCES IN FILE CA (1962 TO DATE)  
 2 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 114:192593  
 REFERENCE 2: 72:3631

L94 ANSWER 26 OF 43 REGISTRY COPYRIGHT 2003 ACS  
 RN 11138-66-2 REGISTRY

CN Xanthan gum (9CI) (CA INDEX NAME)

OTHER NAMES:

CN Actigum CX 9  
 CN ADM 40  
 CN B 1459  
 CN Biopolymer 9702  
 CN Biopolymer XB 23  
 CN Biozan R  
 CN Bisfect XA 200  
 CN Bistop  
 CN Chemicogel  
 CN Echogum  
 CN Echogum F  
 CN Echogum RD  
 CN Echogum SF  
 CN Echogum T  
 CN Ekogum  
 CN Ekogum ketorol  
 CN Enorflo X  
 CN Flocon 1035  
 CN Flocon 4800  
 CN Flocon 4800C  
 CN Flodrill S  
 CN Galaxy XB  
 CN Gums, xanthomonas  
 CN Idvis  
 CN Jungbunzlauer ST  
 CN K 5C151  
 CN K 9C57  
 CN Kelco BT  
 CN Kelflo  
 CN Keltrol  
 CN Keltrol CG  
 CN Keltrol F  
 CN Keltrol RD  
 CN Keltrol SF  
 CN Keltrol T  
 CN Keltrol TF  
 CN Keltrol TF 1000  
 CN Kelzan  
 CN Kelzan 140X  
 CN Kelzan AR  
 CN Kelzan ASX  
 CN Kelzan D  
 CN Kelzan F  
 CN Kelzan M  
 CN Kelzan MF  
 CN Kelzan S  
 CN Kelzan SS 4000  
 CN Kelzan T  
 CN Kelzan XC  
 CN Kelzan XCD

ADDITIONAL NAMES NOT AVAILABLE IN THIS FORMAT - Use FCN, FIDE, or ALL for  
 DISPLAY

DR 12673-42-6, 12771-06-1, 9088-32-8, 54511-23-8, 56592-13-3, 98112-77-7,  
 51811-95-1, 37189-49-4, 37279-85-9, 37332-19-7, 37383-52-1, 80450-59-5,  
 85568-76-9, 82600-55-3, 39393-27-6, 39444-54-7

MF Unspecified

CI PMS, COM, MAN

PCT Manual registration, Polyester, Polyester formed

LC STN Files: AGRICOLA, ANABSTR, AQUIRE, BIOBUSINESS, BIOSIS, BIOTECHNO,  
 CA, CANCERLIT, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS, CHEMLIST, CIN,  
 CSCHEM, DDFU, DRUGU, EMBASE, ENCOMPLIT, ENCOMPLIT2, ENCOMPPAT,



ENCOMPAT2, IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, MRCK\*, MSDS-OHS,  
NAPRALERT, NIOSHTIC, PIRA, PROMT, TOXCENTER, TULSA, USPAT2, USPATFULL,  
VTB

(\*File contains numerically searchable property data)  
Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*  
(\*Enter CHEMLIST File for up-to-date regulatory information)

\*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\*

7112 REFERENCES IN FILE CA (1962 TO DATE)  
243 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
7126 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 138:78512  
REFERENCE 2: 138:78509  
REFERENCE 3: 138:78504  
REFERENCE 4: 138:78503  
REFERENCE 5: 138:78303  
REFERENCE 6: 138:78210  
REFERENCE 7: 138:78187  
REFERENCE 8: 138:77321  
REFERENCE 9: 138:77217  
REFERENCE 10: 138:75726

L94 ANSWER 27 OF 43 REGISTRY COPYRIGHT 2003 ACS  
RN 10043-52-4 REGISTRY  
CN Calcium chloride (CaCl<sub>2</sub>) (9CI) (CA INDEX NAME)  
OTHER CA INDEX NAMES:  
CN Calcium chloride (8CI)  
OTHER NAMES:  
CN Bovikalc  
CN Calcium dichloride  
CN Calcium(2+) chloride  
CN Calcosan  
CN Calmate R  
CN Calol  
CN Calzina oral  
CN Chrysoxal C 4  
CN Daraccel  
CN Dowflake  
CN Liquidow  
CN Peladow  
CN Stopit  
CN U-Ramin MC  
DR 139468-93-2  
MF Ca Cl<sub>2</sub>  
CI COM  
LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, AQUIRE, BIOBUSINESS, BIOSIS,

BIOTECHNO, CA, CABA, CANCERLIT, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS,  
CHEMINFORMRX, CHEMLIST, CIN, CSCHEM, CSNB, DDFU, DETHERM\*, DIOGENES,  
DIPPR\*, DRUGU, EMBASE, ENCOMPLIT, ENCOMPLIT2, ENCOMPAT, ENCOMPAT2,  
GMELIN\*, HSDB\*, IFICDB, IFIPAT, IFIUDB, MEDLINE, MRCK\*, MSDS-OHS,  
NIOSHTIC, PDLCOM\*, PIRA, PROMT, RTECS\*, TOXCENTER, TULSA, USAN, USPAT2,  
USPATFULL, VETU, VTB  
(\*File contains numerically searchable property data)

Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*  
(\*Enter CHEMLIST File for up-to-date regulatory information)

Cl-Ca-Cl

32025 REFERENCES IN FILE CA (1962 TO DATE)  
216 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
32052 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 138:82531

REFERENCE 2: 138:82368

REFERENCE 3: 138:78520

REFERENCE 4: 138:78182

REFERENCE 5: 138:78033

REFERENCE 6: 138:78022

REFERENCE 7: 138:77756

REFERENCE 8: 138:77217

REFERENCE 9: 138:76421

REFERENCE 10: 138:75989

L94 ANSWER 28 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 9050-04-8 REGISTRY

CN Cellulose, carboxymethyl ether, calcium salt (9CI) (CA INDEX NAME)

OTHER NAMES:

CN Ca carboxymethyl cellulose

CN Calcium carboxymethyl cellulose

CN Calcium cellulose glycolate

CN Calcium CM-cellulose

CN Carboxymethyl cellulose calcium salt

CN Carboxymethylcellulose Ca salt

CN Carmellose calcium

CN Cellulose glycolate calcium salt

CN CM-cellulose calcium

CN CM-Cellulose calcium salt

CN ECG 505

CN Hylon 600

DR 9060-76-8, 57406-72-1, 111941-77-6, 52627-96-0

MF C2 H4 O3 . x Ca . x Unspecified

CI COM

PCT Manual registration

LC STN Files: ANABSTR, BIOSIS, BIOTECHNO, CA, CAPLUS, CHEMCATS, CHEMLIST,  
CSCHEM, EMBASE, IFICDB, IFIPAT, IFIUDB, IPA, MSDS-OHS, RTECS\*,  
TOXCENTER, USAN, USPATFULL

(\*File contains numerically searchable property data)

Other Sources: NDSL\*\*, TSCA\*\*

(\*Enter CHEMLIST File for up-to-date regulatory information)

CM 1

CRN 9004-34-6

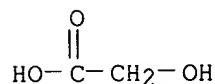
CMF Unspecified

CCI PMS, MAN

\*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\*

CM 2

CRN 79-14-1  
CMF C2 H4 O3



363 REFERENCES IN FILE CA (1962 TO DATE)  
2 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
365 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 138:78444  
REFERENCE 2: 138:66646  
REFERENCE 3: 138:61321  
REFERENCE 4: 137:342154  
REFERENCE 5: 137:342124  
REFERENCE 6: 137:316117  
REFERENCE 7: 137:316064  
REFERENCE 8: 137:268435  
REFERENCE 9: 137:237785  
REFERENCE 10: 137:237763

L94 ANSWER 29 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 9005-70-3 REGISTRY

CN Sorbitan, tri-(9Z)-9-octadecenoate, poly(oxy-1,2-ethanediyl) derivs. (9CI)  
(CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Sorbitan, tri-9-octadecenoate, poly(oxy-1,2-ethanediyl) derivs., (Z,Z,Z)-

CN Sorbitan, trioleate, polyoxyethylene derivs. (8CI)

OTHER NAMES:

CN Alkamuls T 85

CN Crill 12

CN Crillet 45

CN Emsorb 6903

CN Emsorb 6913

CN Ethoxylated sorbitan trioleate

CN Eumulgin STO 80

CN Glycosperse TO 20

CN Montanox 85

CN Newcol 3-85

CN Nikkol TO 30

CN Poly(oxyethylene) sorbitan trioleate

CN Polyethylene glycol sorbitan trioleate

CN Polyoxyethylene sorbitan trioleic acid ester

CN Polysorbate 85

CN Rheodol TW-O 320

CN Rikemal O 852

CN Sinopol 3-85  
 CN Sorbax PTO 20  
 CN Sorbimacrogol trioleate 300  
 CN Sorbitan polyethoxy trioleate  
 CN Sorbitan trioleate polyoxyethylene ether  
 CN Sorbon T 85  
 CN T-MAZ 85  
 CN TO 55  
 CN TO 65  
 CN Toximul SEE 340  
 CN Tween 85  
 CN Witconol AL 69-66  
 DR 9015-60-5, 1340-86-9, 51367-06-7, 111389-05-0  
 MF Unspecified  
 CI PMS, COM, MAN  
 PCT Manual registration  
 LC STN.Files: AGRICOLA, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CAPLUS,  
 CHEMCATS, CHEMLIST, CIN, CSCHEM, EMBASE, IFICDB, IFIPAT, IFIUDB, IPA,  
 MSDS-OHS, PIRA, RTECS\*, TOXCENTER, USAN, USPAT2, USPATFULL  
 (\*File contains numerically searchable property data)  
 Other Sources: DSL\*\*, TSCA\*\*  
 (\*\*Enter CHEMLIST File for up-to-date regulatory information)

\*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\*  
 772 REFERENCES IN FILE CA (1962 TO DATE)  
 8 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
 774 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 138:78450  
 REFERENCE 2: 138:44585  
 REFERENCE 3: 138:40823  
 REFERENCE 4: 138:40671  
 REFERENCE 5: 138:28945  
 REFERENCE 6: 138:26966  
 REFERENCE 7: 138:1959  
 REFERENCE 8: 137:389139  
 REFERENCE 9: 137:365972  
 REFERENCE 10: 137:358152

L94 ANSWER 30 OF 43 REGISTRY COPYRIGHT 2003 ACS  
 RN 9005-64-5 REGISTRY  
 CN Sorbitan, monododecanoate, poly(oxy-1,2-ethanediyl) derivs. (9CI) (CA  
 INDEX NAME)  
 OTHER NAMES:  
 CN Ahco 7596T  
 CN Alkamuls PSML 20  
 CN Alkamuls T 20  
 CN Armotan PML 20  
 CN Atlas G 4280  
 CN Atlas G 7596J  
 CN Atlas G 7596P  
 CN Atmer 110  
 CN Crillet 1  
 CN Disponil SML 120

CN Emasol 1112  
CN Emasol L 130  
CN Emsorb 6915  
CN Ethoxylated sorbitan monolaurate  
CN Ethylene oxide-sorbitan monolaurate adduct  
CN Ethylene oxide-sorbitan monolaurate polymer  
CN Eumulgin SML 15  
CN Eumulgin SML 20  
CN G 1020  
CN G 4280  
CN G 7596J  
CN G 7606J  
CN GL 1  
CN GL 1 (carbohydrate)  
CN Glytanox 1001  
CN Ionet T 20C  
CN Kemotan T 20  
CN Liposorb L 20  
CN LT 221  
CN ML 55F  
CN Montanox 20  
CN Nikkol TL 10  
CN Nissan Nonion LT 204  
CN Nissan Nonion LT 221  
CN Nonion LT 221  
CN Oxyethylated sorbitan monolaurate  
CN Oxysorbic 20  
CN POE sorbitan monolaurate  
CN Poly(ethylene glycol) sorbitan ether monolaurate  
CN Poly(oxyethylene sorbitan laurate)  
CN Poly(oxyethylene)sorbitan ether monolaurate  
CN Poly(oxyethylene)sorbitan monolaurate  
CN Polyethylene glycol sorbitan monolaurate  
CN Polyoxethylene sorbitan monolaurate  
CN Polyoxyethylene sorbitan monododecanoate  
CN Polyoxyethylene Span 20  
CN Polysorbate 20  
CN Polysorbate 21  
CN Polysten 20  
CN Radasurf 7137

ADDITIONAL NAMES NOT AVAILABLE IN THIS FORMAT - Use FCN, FIDE, or ALL for  
DISPLAY

DR 8036-82-6, 9011-30-7, 9015-57-0, 1341-06-6, 122304-31-8, 54174-54-8,  
60318-54-9, 129428-64-4, 62229-28-1, 118955-39-8, 37310-96-6, 93037-36-6,  
194879-92-0

MF Unspecified

CI PMS, COM, MAN

PCT Manual registration, Polyether

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, AQUIRE, BIOBUSINESS, BIOSIS,  
BIOTECHNO, CA, CAPLUS, CASREACT, CHEMCATS, CHEMLIST, CIN, CSCHEM, CSNB,  
DDFU, DRUGU, EMBASE, IFICDB, IFIPAT, IFIUDB, IPA, MSDS-OHS, NIOSHTIC,  
PDLCOM\*, PIRA, PROMT, RTECS\*, TOXCENTER, USAN, USPAT2, USPATFULL  
(\*File contains numerically searchable property data)  
Other Sources: DSL\*\*, TSCA\*\*  
(\*\*Enter CHEMLIST File for up-to-date regulatory information)

\*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\*

5076 REFERENCES IN FILE CA (1962 TO DATE)

25 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

5092 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 138:78459

REFERENCE 2: 138:78333  
 REFERENCE 3: 138:78256  
 REFERENCE 4: 138:78210  
 REFERENCE 5: 138:77217  
 REFERENCE 6: 138:75128  
 REFERENCE 7: 138:75009  
 REFERENCE 8: 138:74052  
 REFERENCE 9: 138:72181  
 REFERENCE 10: 138:72140

L94 ANSWER 31 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 9003-11-6 REGISTRY

CN Oxirane, methyl-, polymer with oxirane (9CI) (CA INDEX NAME)

OTHER NAMES:

CN  $\alpha$ -Hydro- $\omega$ -hydroxy-poly(oxyethylene)-poly(oxypropylene)

CN 1,2-Propanediol polymer with ethylene oxide

CN 333E

CN 50MB-26X

CN 75H380000

CN 75H90000

CN Acclaim 2220N

CN Actcol MF 12

CN Actcol MF 18

CN Actinol P 3035

CN Adeka Carpol MH 150

CN Adeka Carpol MH 500

CN Adeka Carpol PH 2000

CN Adeka CM 294

CN Adeka L 31

CN Adeka PR 3007

CN Adekanol NP 1200

CN Arlatone F 127G

CN Balab 615

CN Berol 370

CN Berol 374

CN Berol TVM 370

CN Bloatguard

CN Breox 50A1000

CN Breox 75W270

CN Breox PAG 50A1000

CN BSP 5000

CN Carpol 2040

CN Carpol 2050

CN CE

CN CF 0802

CN CP 1000

CN CP 1000 (polyoxyalkylene)

CN CP 2000L

CN DE 1

CN DE 1 (demulsifier)

CN Desmophen 7100

CN Dezemulsionat E 96

CN Disfoam CC 222

CN Dissolvan 4411

CN Emkalyx EP 64

CN Emkalyx L 101  
 CN Emulgen PP  
 CN Emulgen PP 150  
 CN Emulgen PP 250  
 CN Emulgen PP 290  
 CN EP 1660  
 CN Epan 420  
 CN Epan 610  
 CN Epan 720

ADDITIONAL NAMES NOT AVAILABLE IN THIS FORMAT - Use FCN, FIDE, or ALL for  
 DISPLAY

AR 53637-25-5  
 DR 452069-47-5, 12676-40-3, 12772-49-5, 9003-12-7, 9009-02-3, 9009-03-4,  
 9009-04-5, 9009-05-6, 9009-06-7, 9010-49-5, 9010-97-3, 9015-66-1,  
 9050-44-6, 9061-69-2, 9067-43-0, 167267-50-7, 168018-54-0, 163032-64-2,  
 163063-49-8, 162627-00-1, 172306-19-3, 53637-72-2, 57971-91-2, 58968-65-3,  
 56730-46-2, 57219-38-2, 57571-70-7, 124057-63-2, 59494-33-6, 59794-22-8,  
 60328-61-2, 64940-96-1, 66746-25-6, 106717-66-2, 50643-24-8, 51312-31-3,  
 51569-27-8, 60976-75-2, 37211-19-1, 37211-20-4, 37211-21-5, 37211-22-6,  
 37211-23-7, 37211-24-8, 37221-18-4, 37265-39-7, 37307-38-3, 37331-16-1,  
 37331-17-2, 37341-81-4, 70213-25-1, 72319-37-0, 73158-62-0, 70644-95-0,  
 71343-56-1, 77448-18-1, 77752-09-1, 76050-76-5, 86249-84-5, 86304-35-0,  
 81180-56-5, 87912-55-8, 91858-59-2, 30600-73-8, 39277-80-0, 39316-56-8,  
 39316-57-9, 39364-13-1, 39387-54-7, 208342-25-0, 232598-91-3, 250780-00-8,  
 254903-86-1, 291775-89-8, 374624-82-5

MF (C3 H6 O . C2 H4 O)x

CI PMS, COM

PCT Polyether, Polyether formed

LC STN Files: AGRICOLA, ANABSTR, AQUIRE, BIOBUSINESS, BIOSIS, BIOTECHNO,  
 CA, CANCERLIT, CAPLUS, CASREACT, CBNB, CHEMLIST, CIN, CSCHEM, DDFU,  
 DIOGENES, DRUGU, EMBASE, ENCOMPLIT, ENCOMPLIT2, ENCOMPPAT, ENCOMPPAT2,  
 IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, MRCK\*, MSDS-OHS, NIOSHTIC, PHAR,  
 PIRA, PROMT, RTECS\*, TOXCENTER, USAN, USPAT2, USPATFULL

(\*File contains numerically searchable property data)

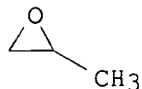
Other Sources: DSL\*\*, TSCA\*\*, WHO

(\*\*Enter CHEMLIST File for up-to-date regulatory information)

CM 1

CRN 75-56-9

CMF C3 H6 O



CM 2

CRN 75-21-8

CMF C2 H4 O



8110 REFERENCES IN FILE CA (1962 TO DATE)  
 2595 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
 8118 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 138:78482  
REFERENCE 2: 138:78474  
REFERENCE 3: 138:78451  
REFERENCE 4: 138:78449  
REFERENCE 5: 138:78444  
REFERENCE 6: 138:78194  
REFERENCE 7: 138:78146  
REFERENCE 8: 138:77327  
REFERENCE 9: 138:75914  
REFERENCE 10: 138:75908

L94 ANSWER 32 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 9000-07-1 REGISTRY

CN Carrageenan (9CI) (CA INDEX NAME)

OTHER NAMES:

CN κλ-Carrageenan  
CN 22: PN: WO0175077 SEQID: 25 claimed sequence  
CN Aubygum x 2  
CN Aubygum X 23  
CN Carrageenan GH  
CN Carrageenan gum  
CN Carrageenan SWG-J  
CN Carrageenin  
CN Carragheen  
CN Carragheenan  
CN EC 4000  
CN FK 6101  
CN FK 6120  
CN Gelcarin GP 37ANF  
CN Gelcarin HWG  
CN Gelloid J  
CN Gelozone  
CN Genugel LC 4  
CN Genugel LC 5  
CN Genugel MG 11  
CN Genugel RLV  
CN Genuvisco J  
CN Genuvisco TPH-1  
CN Gum carrageenan  
CN Gum chon 2  
CN Gum chond  
CN Inagel E 150  
CN Isagel RG 300  
CN LSS 1  
CN ME 913  
CN Newgelin LB 4  
CN Norsk gelatan  
CN Pellugel  
CN Pencogel  
CN Satiagel NP 5B  
CN Sea-Pi Gum FA  
CN Seagel GH  
CN Seagel Pet  
CN SeaKem carrageenin



CN SeaKem CM 611  
 CN Sherex IC 109  
 CN Soa Ace WX 138  
 CN Soa Ace WX 165  
 CN Soageena MM 501  
 CN Soageena MW 351  
 CN Soageena WX 560  
 CN T 307 (gellant)  
 CN Takaragen G 50  
 CN Takaragen L  
 CN TIC Pretested Colloid 775

ADDITIONAL NAMES NOT AVAILABLE IN THIS FORMAT - Use FCN, FIDE, or ALL for  
 DISPLAY

DR 8040-42-4, 9000-13-9, 9000-27-5, 78005-48-8

MF Unspecified

CI PMS, COM, MAN

PCT Manual registration, Polyother, Polyother only

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, BIOBUSINESS, BIOSIS, BIOTECHNO,  
 CA, CABA, CANCERLIT, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS, CHEMLIST,  
 CIN, CSCHEM, CSNB, DDFU, DRUGU, EMBASE, ENCOMPLIT, ENCOMPLIT2,  
 ENCOMPPAT, ENCOMPPAT2, HSDB\*, IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE,  
 MRCK\*, MSDS-OHS, NAPRALERT, NIOSHTIC, PIRA, PROMT, RTECS\*, TOXCENTER,  
 USAN, USPAT2, USPATFULL

(\*File contains numerically searchable property data)

Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*

(\*\*Enter CHEMLIST File for up-to-date regulatory information)

\*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\*

4174 REFERENCES IN FILE CA (1962 TO DATE)

125 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

4183 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 138:78509

REFERENCE 2: 138:78210

REFERENCE 3: 138:78187

REFERENCE 4: 138:77609

REFERENCE 5: 138:72585

REFERENCE 6: 138:72314

REFERENCE 7: 138:72087

REFERENCE 8: 138:72069

REFERENCE 9: 138:66907

REFERENCE 10: 138:66690

L94 ANSWER 33 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 7487-88-9 REGISTRY

CN Sulfuric acid magnesium salt (1:1) (8CI, 9CI) (CA INDEX NAME)

OTHER NAMES:

CN Epsom salt

CN Epsom salts

CN Magnesium sulfate

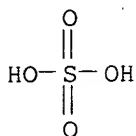
CN Magnesium sulfate (1:1)

CN Magnesium sulphate

CN OT-S

CN OT-S (drying agent)

CN Sulfuric acid magnesium salt  
 CN Sulfuric acid, magnesium salt (1:1)  
 CN Tomix OT  
 AR 18939-43-0  
 DR 139939-75-6  
 MF H2 O4 S . Mg  
 CI COM  
 LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, AQUIRE, BIOBUSINESS, BIOSIS,  
 BIOTECHNO, CA, CABA, CANCERLIT, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS,  
 CHEMINFORMRX, CHEMLIST, CIN, CSCHEM, CSNB, DDFU, DETHERM\*, DIOGENES,  
 DIPPR\*, DRUGU, EMBASE, ENCOMPLIT, ENCOMPLIT2, ENCOMPPAT, ENCOMPPAT2,  
 GMELIN\*, HSDB\*, IFICDB, IFIPAT, IFIUDB, MEDLINE, MRCK\*, MSDS-OHS,  
 NIOSHTIC, PDLCOM\*, PIRA, PROMT, RTECS\*, TOXCENTER, TULSA, USAN, USPAT2,  
 USPATFULL, VETU, VTB  
 (\*File contains numerically searchable property data)  
 Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*  
 (\*\*Enter CHEMLIST File for up-to-date regulatory information)  
 CRN (7664-93-9)



● Mg

12483 REFERENCES IN FILE CA (1962 TO DATE)  
 87 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
 12498 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 138:81747  
 REFERENCE 2: 138:77373  
 REFERENCE 3: 138:76301  
 REFERENCE 4: 138:74989  
 REFERENCE 5: 138:73378  
 REFERENCE 6: 138:72644  
 REFERENCE 7: 138:72336  
 REFERENCE 8: 138:72185  
 REFERENCE 9: 138:72011  
 REFERENCE 10: 138:72006

L94 ANSWER 34 OF 43 REGISTRY COPYRIGHT 2003 ACS  
 RN 7440-70-2 REGISTRY  
 CN Calcium (8CI, 9CI) (CA INDEX NAME)  
 OTHER NAMES:  
 CN Atomic calcium  
 CN Blood-coagulation factor IV

CN Calcium atom  
 CN Calcium element  
 CN Praval  
 DR 8047-59-4  
 MF Ca  
 CI COM  
 LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, AQUIRE, BIOBUSINESS, BIOSIS,  
 BIOTECHNO, CA, CABA, CANCERLIT, CAOLD, CAPLUS, CASREACT, CBNB, CEN,  
 CHEMCATS, CHEMINFORMRX, CHEMLIST, CIN, CSCHEM, CSNB, DDFU, DETHERM\*,  
 DIOGENES, DIPPR\*, DRUGU, EMBASE, ENCOMPLIT, ENCOMPLIT2, ENCOMPPAT,  
 ENCOMPPAT2, HSDB\*, IFICDB, IFIPAT, IFIADB, IPA, MEDLINE, MRCK\*,  
 MSDS-OHS, NAPRALERT, NIOSHTIC, PHARMASEARCH, PIRA, PROMT, TOXCENTER,  
 TULSA, ULIDAT, USPAT2, USPATFULL, VETU, VTB  
 (\*File contains numerically searchable property data)  
 Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*  
 (\*\*Enter CHEMLIST File for up-to-date regulatory information)

Ca

305816 REFERENCES IN FILE CA (1962 TO DATE)  
 6524 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
 305925 REFERENCES IN FILE CAPLUS (1962 TO DATE)  
 1 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

REFERENCE 1: 138:82583  
 REFERENCE 2: 138:82570  
 REFERENCE 3: 138:82568  
 REFERENCE 4: 138:82519  
 REFERENCE 5: 138:82515  
 REFERENCE 6: 138:82488  
 REFERENCE 7: 138:81750  
 REFERENCE 8: 138:80919  
 REFERENCE 9: 138:80778  
 REFERENCE 10: 138:80777

L94 ANSWER 35 OF 43 REGISTRY COPYRIGHT 2003 ACS  
 RN 7440-66-6 REGISTRY  
 CN Zinc (7CI, 8CI, 9CI) (CA INDEX NAME)  
 OTHER NAMES:  
 CN AN 325  
 CN Asarco L 15  
 CN Blue powder  
 CN Ecka 4  
 CN F 1000  
 CN F 1000 (metal)  
 CN F 1500T  
 CN F 2000  
 CN F 2000 (metal)  
 CN LS 2  
 CN LS 2 (element)  
 CN LS 4  
 CN LS 5

CN LS 5 (metal)  
 CN MCS  
 CN MCS (metal)  
 CN NC-Zinc  
 CN Rheinzink  
 CN UF  
 CN UF (metal)  
 CN VM 4P16  
 CN Zinc Dust 3  
 DR 12793-53-2, 195161-85-4, 199281-21-5, 298688-49-0  
 MF Zn  
 CI COM  
 LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, AQUIRE, BIOBUSINESS, BIOSIS,  
 BIOTECHNO, CA, CABA, CANCERLIT, CAOLD, CAPLUS, CASREACT, CBNB, CEN,  
 CHEMCATS, CHEMINFORMRX, CHEMLIST, CHEMSAFE, CIN, CSCHEM, CSNB, DDFU,  
 DETHERM\*, DIOGENES, DIPPR\*, DRUGU, EMBASE, ENCOMPLIT, ENCOMPLIT2,  
 ENCOMPPAT, ENCOMPPAT2, HSDB\*, IFICDB, IFIPAT, IFIADB, IPA, MEDLINE,  
 MRCK\*, MSDS-OHS, NAPRALERT, NIOSHTIC, PDLCOM\*, PHARMASEARCH, PIRA,  
 PROMT, RTECS\*, TOXCENTER, TULSA, ULIDAT, USPAT2, USPATFULL, VETU, VTB  
 (\*File contains numerically searchable property data)  
 Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*  
 (\*\*Enter CHEMLIST File for up-to-date regulatory information)

Zn

224704 REFERENCES IN FILE CA (1962 TO DATE)  
 11496 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
 224822 REFERENCES IN FILE CAPLUS (1962 TO DATE)  
 1 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

REFERENCE 1: 138:82601  
 REFERENCE 2: 138:82568  
 REFERENCE 3: 138:82547  
 REFERENCE 4: 138:82543  
 REFERENCE 5: 138:82525  
 REFERENCE 6: 138:82519  
 REFERENCE 7: 138:82517  
 REFERENCE 8: 138:82488  
 REFERENCE 9: 138:82487  
 REFERENCE 10: 138:82303

L94 ANSWER 36 OF 43 REGISTRY COPYRIGHT 2003 ACS  
 RN 7440-50-8 REGISTRY  
 CN Copper (7CI, 8CI, 9CI) (CA INDEX NAME)  
 OTHER NAMES:  
 CN 100RXH  
 CN 1100T  
 CN 115A  
 CN 1721 Gold  
 CN 200RL  
 CN 22BB400  
 CN 2L3GT

CN 3EC  
CN 3EC-HTE  
CN 3EC-III  
CN 3EC-VLP  
CN 3EC3  
CN 3L Fire  
CN Allbri Natural Copper  
CN Arwood copper  
CN BHN 02T  
CN BHY 02B-T  
CN BHY 13HT  
CN BHY 13T  
CN BHY 22B-T  
CN BPF 18  
CN BSH  
CN BSH (metal)  
CN C 100  
CN C 100 (metal)  
CN C.I. 77400  
CN C.I. Pigment Metal 2  
CN CDX  
CN CDX (metal)  
CN CE 1100  
CN CE 1110  
CN CE 115  
CN CE 15  
CN CE 25  
CN CE 7  
CN CE 7 (metal)  
CN CE 8A  
CN CF 78  
CN CF-T 8  
CN Copper element  
CN Copper Powder  
CN CS-F 150E  
CN CT 315E  
CN CU 112  
CN Cu-At-W 250  
CN CU-FN 10  
CN Cu-HWQ  
CN CuEP  
CN CuEPP  
CN CuLox 6010

ADDITIONAL NAMES NOT AVAILABLE IN THIS FORMAT - Use FCN, FIDE, or ALL for  
DISPLAY

DR 133353-46-5, 133353-47-6, 65555-90-0, 72514-83-1, 195161-80-9

MF Cu

CI COM

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, AQUIRE, BIOBUSINESS, BIOSIS,  
BIOTECHNO, CA, CABA, CANCERLIT, CAOLD, CAPLUS, CASREACT, CBNB, CEN,  
CHEMCATS, CHEMINFORMRX, CHEMLIST, CHEMSAFE, CIN, CSCHEM, CSNB, DDFU,  
DETERM\*, DIOGENES, DRUGU, EMBASE, ENCOMPLIT, ENCOMPLIT2, ENCOMPPAT,  
ENCOMPPAT2, HSDB\*, IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, MRCK\*,  
MSDS-OHS, NAPRALERT, NIOSHTIC, PIRA, PROMT, RTECS\*, TOXCENTER, TULSA,  
ULIDAT, USPAT2, USPATFULL, VETU, VTB

(\*File contains numerically searchable property data)

Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*

(\*\*Enter CHEMLIST File for up-to-date regulatory information)

401834 REFERENCES IN FILE CA (1962 TO DATE)  
20914 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
402077 REFERENCES IN FILE CAPLUS (1962 TO DATE)  
2 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

REFERENCE 1: 138:82644  
REFERENCE 2: 138:82585  
REFERENCE 3: 138:82584  
REFERENCE 4: 138:82570  
REFERENCE 5: 138:82567  
REFERENCE 6: 138:82547  
REFERENCE 7: 138:82521  
REFERENCE 8: 138:82519  
REFERENCE 9: 138:82517  
REFERENCE 10: 138:82488

L94 ANSWER 37 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 7439-96-5 REGISTRY

CN Manganese (8CI, 9CI) (CA INDEX NAME)

OTHER NAMES:

CN Colloidal manganese

CN Cutaval

CN JIS-G 1213

CN Manganese element

CN Manganese fulleride (MnC20)

CN Manganese-55

DR 8031-40-1, 8075-39-6, 17375-02-9, 39303-06-5, 195161-78-5

MF Mn

CI COM

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, AQUIRE, BIOBUSINESS, BIOSIS,  
BIOTECHNO, CA, CABA, CANCERLIT, CAOLD, CAPLUS, CASREACT, CBNB, CEN,  
CHEMCATS, CHEMLIST, CHEMSAFE, CIN, CSCHEM, CSNB, DDFU, DETHERM\*,  
DIOGENES, DRUGU, EMBASE, ENCOMPLIT, ENCOMPLIT2, ENCOMPPAT, ENCOMPPAT2,  
HSDB\*, IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, MRCK\*, MSDS-OHS, NIOSHTIC,  
PIRA, PROMT, RTECS\*, TOXCENTER, TULSA, ULIDAT, USPAT2, USPATFULL, VETU,  
VTB

(\*File contains numerically searchable property data)

Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*

(\*\*Enter CHEMLIST File for up-to-date regulatory information)

Mn

141579 REFERENCES IN FILE CA (1962 TO DATE)  
7014 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
141690 REFERENCES IN FILE CAPLUS (1962 TO DATE)  
1 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

REFERENCE 1: 138:82601  
REFERENCE 2: 138:82585  
REFERENCE 3: 138:82568

REFERENCE 4: 138:82562  
 REFERENCE 5: 138:82519  
 REFERENCE 6: 138:82517  
 REFERENCE 7: 138:82488  
 REFERENCE 8: 138:82466  
 REFERENCE 9: 138:82136  
 REFERENCE 10: 138:82123

L94 ANSWER 38 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 7439-95-4 REGISTRY

CN Magnesium (8CI, 9CI) (CA INDEX NAME)

OTHER NAMES:

CN JIS 1

CN Magnesium element

CN PK 31

CN PK 31 (magnesium)

CN Rieke's active magnesium

DR 14147-08-1, 67208-78-0, 199281-20-4, 298688-48-9

MF Mg

CI COM

LC STN Files: ANABSTR, AQUIRE, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CABA, CANCERLIT, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS, CHEMINFORMRX, CHEMLIST, CIN, CSCHM, CSNB, DDFU, DETHERM\*, DRUGU, EMBASE, ENCOMPLIT, ENCOMPLIT2, ENCOMPPAT, ENCOMPPAT2, HSDB\*, IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, MRCK\*, MSDS-OHS, NAPRALERT, NIOSHTIC, TOXCENTER, ULIDAT, USPAT2, USPATFULL, VETU, VTB

(\*File contains numerically searchable property data)

Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*

(\*\*Enter CHEMLIST File for up-to-date regulatory information)

Mg

165199 REFERENCES IN FILE CA (1962 TO DATE)

6224 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

165330 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 138:82570  
 REFERENCE 2: 138:82568  
 REFERENCE 3: 138:82519  
 REFERENCE 4: 138:82488  
 REFERENCE 5: 138:82313  
 REFERENCE 6: 138:82302  
 REFERENCE 7: 138:82042  
 REFERENCE 8: 138:81947  
 REFERENCE 9: 138:81750

REFERENCE 10: 138:81743

L94 ANSWER 39 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 7439-93-2 REGISTRY

CN Lithium (7CI, 8CI, 9CI) (CA INDEX NAME)

OTHER NAMES:

CN Lithium atom

CN Lithium element

MF Li

CI COM

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, AQUIRE, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CABA, CANCERLIT, CAOLD, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS, CHEMINFORMRX, CHEMLIST, CIN, CSCHEM, CSNB, DDFU, DETHERM+, DIPPR\*, DRUGU, EMBASE, ENCOMPLIT, ENCOMPLIT2, ENCOMPPAT, ENCOMPPAT2, HSDB\*, IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, MRCK\*, MSDS-OHS, NIOSHTIC, PIRA, PROMT, RTECS\*, TOXCENTER, TULSA, ULIDAT, USPAT2, USPATFULL, VETU, VTB

(\*File contains numerically searchable property data)

Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*

(\*\*Enter CHEMLIST File for up-to-date regulatory information)

Li

64251 REFERENCES IN FILE CA (1962 TO DATE)

5310 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

64293 REFERENCES IN FILE CAPLUS (1962 TO DATE)

5 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

REFERENCE 1: 138:82519

REFERENCE 2: 138:82370

REFERENCE 3: 138:81743

REFERENCE 4: 138:81499

REFERENCE 5: 138:81141

REFERENCE 6: 138:80391

REFERENCE 7: 138:80357

REFERENCE 8: 138:80350

REFERENCE 9: 138:80335

REFERENCE 10: 138:80155

L94 ANSWER 40 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 7439-89-6 REGISTRY

CN Iron (7CI, 8CI, 9CI) (CA INDEX NAME)

OTHER NAMES:

CN 300A

CN 3ZhP

CN A 227

CN Ancor B

CN Ancor EN 80/150

CN AQ 80

CN Armco iron

CN Atomel 300M200

CN Atomel 500M



CN Atomet 28  
 CN Atomet 95  
 CN Atomiron 44MR  
 CN Atomiron 5M  
 CN Atomiron AFP 25  
 CN Atomiron AFP 5  
 CN ATW 230  
 CN ATW 432  
 CN BASF-EW  
 CN Carbon 0.17, iron 99.83 (atomic)  
 CN Carbonyl iron  
 CN Copy Powder CS 105-175  
 CN DH  
 CN Diseases (animal), iron overload  
 CN Diseases, iron overload  
 CN DSP 1000  
 CN DSP 128B  
 CN DSP 135  
 CN DSP 135C  
 CN DSP 138  
 CN EF 1000  
 CN EF 250  
 CN EFV  
 CN EFV 200/300  
 CN EFV 250  
 CN EFV 250/400  
 CN EO 5A  
 CN F 60  
 CN F 60 (metal)  
 CN Ferrovac E  
 CN FT 3  
 CN FT 3 (element)  
 CN GS 6  
 CN HF 2  
 CN HF 2 (element)  
 CN HL (iron)  
 CN Hoeganaes ATW 230  
 CN Hoeganaes EH  
 CN HQ  
 CN HQ (metal)  
 CN HS (iron)

ADDITIONAL NAMES NOT AVAILABLE IN THIS FORMAT - Use FCN, FIDE, or ALL for  
DISPLAY

DR 8011-79-8, 8053-60-9, 129048-51-7, 73135-38-3, 70884-35-4, 39344-71-3,  
190454-13-8, 195161-83-2, 199281-22-6, 443783-52-6

MF Fe

CI COM

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, AQUIRE, BIOBUSINESS, BIOSIS,  
BIOTECHNO, CA, CABA, CANCERLIT, CAOLD, CAPLUS, CASREACT, CBNB, CEN,  
CHEMCATS, CHEMINFORMRX, CHEMLIST, CIN, CSCHEM, CSNB, DDFU, DETHERM\*,  
DIOGENES, DIPPR\*, DRUGU, EMBASE, ENCOMPLIT, ENCOMPLIT2, ENCOMPPAT,  
ENCOMPPAT2, HSDB\*, IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, MRCK\*,  
MSDS-OHS, NIOSHTIC, PHARMASEARCH, PIRA, PROMT, RTECS\*, TOXCENTER, TULSA,  
ULIDAT, USPAT2, USPATFULL, VETU, VTB

(\*File contains numerically searchable property data)

Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*

(\*\*Enter CHEMLIST File for up-to-date regulatory information)

Fe

17891 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
315740 REFERENCES IN FILE CAPLUS (1962 TO DATE)  
1 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

REFERENCE 1: 138:82585  
REFERENCE 2: 138:82580  
REFERENCE 3: 138:82577  
REFERENCE 4: 138:82573  
REFERENCE 5: 138:82562  
REFERENCE 6: 138:82549  
REFERENCE 7: 138:82537  
REFERENCE 8: 138:82528  
REFERENCE 9: 138:82525  
REFERENCE 10: 138:82522

L94 ANSWER 41 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 7429-90-5 REGISTRY

CN Aluminum (8CI, 9CI) (CA INDEX NAME)

OTHER NAMES:

CN 0670TS  
CN 0900X  
CN 1001M  
CN 102B  
CN 102C  
CN 1100H24  
CN 1100P-H18  
CN 13T  
CN 350D  
CN 350F  
CN 40XD  
CN 5422NS  
CN 5654NS  
CN 5N  
CN 5XD  
CN 716ON  
CN 725EA  
CN 725N  
CN 7620NS  
CN 7640NS  
CN 8011A  
CN 97-4071RE  
CN A 1-18000  
CN A 5052H34  
CN A 6063S  
CN A 95  
CN A 95 (metal)  
CN A 99  
CN A 99 (metal)  
CN A 999  
CN A 999V  
CN A 99N  
CN AA 15  
CN AB 1003  
CN AC 0460

CN AC 1000  
 CN AC 1000 (metal)  
 CN AC 1003  
 CN AC 2500  
 CN AC 5000  
 CN AC 5005  
 CN AIH 30H  
 CN AIH 30H-0  
 CN Aisin Metal Fiber  
 CN Al 050P-H24  
 CN Al 18000  
 CN AL-AT 250  
 CN Al-At 500F  
 CN Albo F  
 CN ALC Fine

ADDITIONAL NAMES NOT AVAILABLE IN THIS FORMAT - Use FCN, FIDE, or ALL for  
 DISPLAY

DR. 12766-45-9, 113962-66-6, 37202-64-5, 80341-19-1, 91728-14-2, 39302-71-1,  
 39332-62-2, 182260-45-3, 185464-37-3, 257888-99-6, 298688-47-8

MF Al

CI COM

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, AQUIRE, BIOBUSINESS, BIOSIS,  
 BIOTECHNO, CA, CABA, CANCERLIT, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS,  
 CHEMINFORMRX, CHEMLIST, CHEMSAFE, CIN, CSCHEM, CSNB, DDFU, DETHERM\*,  
 DIOGENES, DIPPR\*, DRUGU, EMBASE, ENCOMPLIT, ENCOMPLIT2, ENCOMPPAT,  
 ENCOMPPAT2, HSDB\*, IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, MRCK\*,  
 MSDS-OHS, NIOSHTIC, PIRA, PROMT, RTECS\*, TOXCENTER, TULSA, ULIDAT,  
 USPAT2, USPATFULL, VTB

(\*File contains numerically searchable property data)

Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*

(\*\*Enter CHEMLIST File for up-to-date regulatory information)

Al

295751 REFERENCES IN FILE CA (1962 TO DATE)  
 8839 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA  
 295866 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 138:82571

REFERENCE 2: 138:82554

REFERENCE 3: 138:82490

REFERENCE 4: 138:82461

REFERENCE 5: 138:82313

REFERENCE 6: 138:82311

REFERENCE 7: 138:82308

REFERENCE 8: 138:82302

REFERENCE 9: 138:82136

REFERENCE 10: 138:82130

L94 ANSWER 42 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 1492-18-8 REGISTRY

CN L-Glutamic acid, N-[4-[[ (2-amino-5-formyl-1,4,5,6,7,8-hexahydro-4-oxo-6-

pteridiny]methyl]amino]benzoyl]-, calcium salt (1:1) (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Glutamic acid, N-[p-[(2-amino-5-formyl-5,6,7,8-tetrahydro-4-hydroxy-6-pteridiny]methyl]amino]benzoyl]-, calcium salt (1:1), L- (8CI)

OTHER NAMES:

CN Calcium 5-formyltetrahydrofolate

CN Calcium folinate

CN Calcium L-folinate

CN Calcium leucovorin

CN Folinic acid calcium salt

CN Lederfoline

CN Leucovorin calcium

CN Leucovorin calcium salt

CN Rescuvinol

FS STEREOSEARCH

DR 6035-86-5, 6209-45-6, 30771-29-0

MF C20 H23 N7 O7 . Ca

CI COM

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, BEILSTEIN\*, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CAOLD, CAPLUS, CBNB, CHEMCATS, CHEMLIST, CIN, CSCHEM, DDFU, DIOGENES, DRUGU, EMBASE, HSDB\*, MSDS-OHS, PHARMASEARCH, PROMT, RTECS\*, TOXCENTER, USAN, USPATFULL

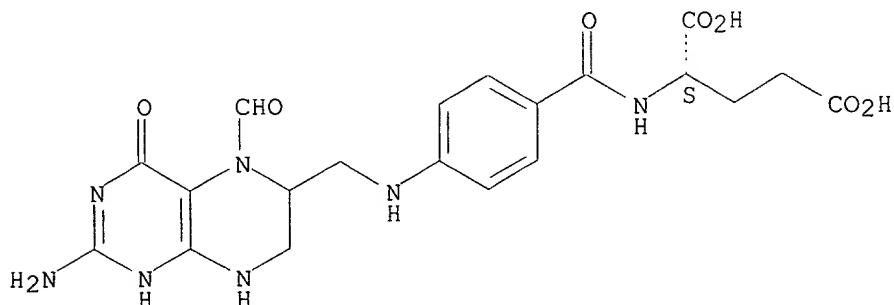
(\*File contains numerically searchable property data)

Other Sources: EINECS\*\*, WHO

(\*\*Enter CHEMLIST File for up-to-date regulatory information)

CRN (58-05-9)

Absolute stereochemistry.



● Ca

131 REFERENCES IN FILE CA (1962 TO DATE)

3 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

131 REFERENCES IN FILE CAPLUS (1962 TO DATE)

1 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

REFERENCE 1: 138:78473

REFERENCE 2: 138:61373

REFERENCE 3: 138:11274

REFERENCE 4: 137:362991

REFERENCE 5: 137:320299

REFERENCE 6: 137:257635

REFERENCE 7: 137:195570

REFERENCE 8: 137:103549

REFERENCE 9: 137:57367

REFERENCE 10: 137:15155

L94 ANSWER 43 OF 43 REGISTRY COPYRIGHT 2003 ACS

RN 299-28-5 REGISTRY

CN D-Gluconic acid, calcium salt (2:1) (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Gluconic acid, calcium salt (2:1), D- (8CI)

OTHER NAMES:

CN Biocal

CN Calcicol

CN Calciofon

CN Calcipur

CN Calcium D-gluconate

CN Calcium gluconate

CN Calcium hexagluconate

CN Calglucol

CN Calglucon

CN Dragocal

CN Ebucin

CN Glucal

CN Glucobiogen

CN Kalpren

CN Novocal

FS STEREOSEARCH

DR 18016-24-5, 3414-35-5

MF C6 H12 O7 . 1/2 Ca

CI COM

LC STN Files: ADISNEWS, AGRICOLA, ANABSTR, BEILSTEIN\*, BIOBUSINESS, BIOSIS, BIOTECHNO, CA, CANCERLIT, CAOLD, CAPLUS, CASREACT, CBNB, CEN, CHEMCATS, CHEMLIST, CIN, CSCHEM, CSNB, DDFU, DETHERM\*, DIOGENES, DRUGU, EMBASE, HSDB\*, IFICDB, IFIPAT, IFIUDB, IPA, MEDLINE, MRCK\*, MSDS-OHS, NAPRALERT, NIOSHTIC, PDLCOM\*, PIRA, PROMT, RTECS\*, SPECINFO, TOXCENTER, TULSA, USAN, USPAT2, USPATFULL, VETU, VTB

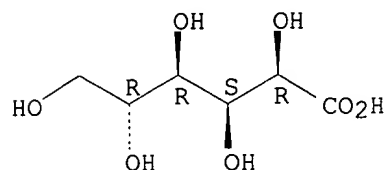
(\*File contains numerically searchable property data)

Other Sources: DSL\*\*, EINECS\*\*, TSCA\*\*

(\*\*Enter CHEMLIST File for up-to-date regulatory information)

CRN (526-95-4)

Absolute stereochemistry.



● 1/2 Ca

1529 REFERENCES IN FILE CA (1962 TO DATE)

10 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

1529 REFERENCES IN FILE CAPLUS (1962 TO DATE)

7 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

lucas 09 / 888114

REFERENCE	1:	138:78539
REFERENCE	2:	138:61315
REFERENCE	3:	138:33365
REFERENCE	4:	138:29206
REFERENCE	5:	138:10749
REFERENCE	6:	138:8350
REFERENCE	7:	138:8313
REFERENCE	8:	138:412
REFERENCE	9:	137:389223
REFERENCE	10:	137:389146